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(57) Abstract

Arrays of polynucleotide spots and kits comprising the same, as well as methods for their preparation and use are provided. The subject arrays include a plurality of polynucleotide spots stably associated with the surface of a solid support. At least a portion of the polynucleotide spots comprises a polynucleotide probe composition that is made up of unique polynucleotides, where all of the unique polynucleotides of the array correspond to a common type of gene. Also provided are sets of a representational number of gene specific primers suitable for use in generating target nucleic acid for use with the subject arrays. The subject arrays find use in hybridization assays, particularly in assays for the identification of differential gene expression patterns among two or more different types of cells.

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NUCLEIC ACID ARRAYS

CROSS-REFERENCE TO RELATED APPLICATIONS

This application is a continuation-in-part of application serial no. 08/859,998 filed on May 21, 1997 and application serial no. 09/053,375 filed on March 31, 1998, the disclosures of which are herein incorporated by reference.

INTRODUCTION

Technical Field

The field of this invention is biopolymeric arrays.

Background of the Invention

"Biochips" or arrays of binding agents, such as oligonucleotides and peptides, have become an increasingly important tool in the biotechnology industry and related fields.

These binding agent arrays, in which a plurality of binding agents are deposited onto a solid support surface in the form of an array or pattern, find use in a variety of applications, including drug screening, nucleic acid sequencing, mutation analysis, and the like. One important use of biochips is in the analysis of differential gene expression, where the expression of genes in different cells, normally a cell of interest and a control, is compared and any discrepancies in expression are identified. In such assays, the presence of discrepancies indicates a difference in the classes of genes expressed in the cells being compared.

In methods of differential gene expression, arrays find use by serving as a substrate to which is bound polynucleotide "probe" fragments. One then obtains "targets" from

analogous cells, tissues or organs of a healthy and diseased organism. The targets are then hybridized to the immobilized set of polynucleotide "probe" fragments. Differences between the resultant hybridization patterns are then detected and related to differences in gene expression in the two sources.

A variety of different array technologies have been developed in order to meet the growing need of the biotechnology industry, as evidenced by the extensive number of patents and references listed in the relevant literature section below.

Despite the wide variety of array technologies currently in preparation or available on the market, there is a continued need to identify new array devices to meet the needs of specific applications. Of particular interest would be the development of an array capable of providing high throughput analysis of differential gene expression.

Relevant Literature

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Patents and patent applications describing arrays of biopolymeric compounds and methods for their fabrication include: 5,242,974; 5,384,261; 5,405,783; 5,412,087; 5,424,186; 5,429,807; 5,436,327; 5,445,934; 5,472,672; 5,527,681; 5,529,756; 5,545,531; 5,554,501; 5,556,752; 5,561,071; 5,599,895; 5,624,711; 5,639,603; 5,658,734; WO 93/17126; WO 95/11995; WO 95/35505; EP 742 287; and EP 799 897.

Patents and patent application describing methods of using arrays in various applications include: 5,143,854; 5,288,644; 5,324,633; 5,432,049; 5,470,710; 5,492,806; 5,503,980; 5,510,270; 5,525,464; 5,547,839; 5,580,732; 5,661,028; WO 95/21265; WO 96/31622; WO 97/10365; WO 97/27317; EP 373 203; and EP 785 280.

Other references of interest include: Atlas Human cDNA Expression Array I (April 1997) CLONTECHniques XII: 4-7; Lockhart et al., Nature Biotechnology (1996) 14: 1675-1680; Shena et al., Science (1995) 270: 467-470; Schena et al., Proc. Nat'l Acad. Sci. USA (1996)93:10614-10619; Shalon et al., Genome Res. (1996) 6: 639-645; Milosavljevic et al., Genome Res. (1996) 6:132-141; Nguyen et al., Genomics (1995)29: 207-216; Piétu et al., Genome Res. (1996) 6: 492-503; Zhao et al., Gene (1995) 166:207-213; Chalifour et al., Anal. Biochem. (1994) 216:299-304; Heller et al., Proc. Nat'l Acad. Sci. USA (1997) 94: 2150-2155; and Schena, M., BioAssays (1996) 18: 427-431.

SUMMARY OF THE INVENTION

Arrays of polynucleotide spots stably associated with the surface of a solid support and kits comprising the same, as well as methods for their preparation and use in hybridization assays, are provided. The subject arrays comprise a plurality of polynucleotide spots, wherein each different polynucleotide spot is made up of a polynucleotide probe composition and at least a portion of the polynucleotide probe compositions are made up of unique polynucleotides. The arrays are further characterized in that all of the unique polynucleotides on the array correspond to the same type of gene. The subject arrays find particular use in differential gene expression analysis. Also provided are sets of a representational number of gene specific primers useful in generating target nucleic acids for use with the subject arrays in hybridization assays.

BRIEF DESCRIPTION OF THE FIGURES

Fig. 1 provides a representation of an array according to the subject invention.

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DEFINITIONS

The term "nucleic acid" as used herein means a polymer composed of nucleotides, e.g. deoxyribonucleotides or ribonucleotides.

The terms "ribonucleic acid" and "RNA" as used herein mean a polymer composed of ribonucleotides.

The terms "deoxyribonucleic acid" and "DNA" as used herein mean a polymer composed of deoxyribonucleotides.

The term "oligonucleotide" as used herein denotes single stranded nucleotide multimers of from about 10 to 100 nucleotides in length.

The term "polynucleotide" as used herein refers to single or double stranded polymer composed of nucleotide monomers of greater than about 120 nucleotides in length up to about 1000 nucleotides in length.

The term "array type" refers to the type of gene represented on the array by the unique polynucleotides, where the type of gene that is represented on the array is dependent on the intended purpose of the array, e.g. to monitor expression of key human genes, to monitor expression of known oncogenes, etc, i.e. the use for which the array is designed. As such, all of the unique polynucleotides on a given array correspond to the same type or

category or group of genes. Genes are considered to be of the same type if they share some common linking characteristics, such as: species of origin, e.g. human, mouse, rat, etc.; tissue or cell type of origin, e.g. muscle, neural, dermal, organ, etc.; disease state, e.g. cancer; functions, e.g. protein kinases, tumor supressors and the like, participation in the same normal biological process, e.g. apoptosis, signal transduction, cell cycle regulation, proliferation, differentiation etc.; and the like. For example, one array type that is provided below is a "cancer array" in which each of the "unique" polynucleotide probes correspond to a gene associated with a cancer disease state. Likewise, a "human array" may be an array of polynucleotides corresponding to unique tightly regulated human genes. Similarly, an "apoptosis array" may be an array type in which the polynucleotides correspond to unique genes associated with apoptosis.

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The "unique" polynucleotide sequences associated with each type of array of the present invention are sequences which are distinctive or different with respect to every other polynucleotide sequence on the array and correspond to the same type of gene, as defined above. For example, in a cancer array, each unique polynucleotide has a sequence that is not homologous to any other known cancer associated sequence. Moreover, each polynucleotide sequence on the array is statistically chosen to ensure that the probability of homology to any sequence of that type is very low. Morever, in the cancer array embodiment, all sequences are statistically chosen to insure that the probability of homology to any other sequence associated with cancer or of human origin is very low. An important feature of the individual polynucleotide probe compositions of the subject arrays is that they are only a fragment of the entire cDNA of the gene to which they correspond. In other words, for each gene represented on the array, the entire cDNA sequence the gene is not represented on the array. Instead, the sequence of only a portion or fragment of the entire cDNA is represented on the array by this unique polynucleotide.

The term "polynucleotide probe composition" refers to the nucleic acid composition that makes up each of the spots on the array. Thus, the term "polynucleotide probe composition" includes nucleic acid compositions of unique polynucleotides and control or calibrating polynucleotides (e.g. polynucleotides corresponding to housekeeping genes). The polynucleotide compositions are made up of single stranded polynucleotides (i.e. polynucleotides that are not hybridized to each other), where all of the polynucleotides in the probe composition may be identical to each other or there may be two different

polynucleotides (polynucleotides of different nucleotide sequence) in each probe composition, where the two different polynucleotides are complementary to each other.

The term "gene specific primer" means a polynucleotide of sufficient length to specifically hybridize to a distinct nucleic acid member of the sample, e.g. RNA or cDNA, where the length of the gene specific primers will usually be at least 8 nt, more usually at least 20 nt and may be as long as 25 nt or longer, but will usually not exceed 50 nt. The gene specific primers of the subject invention are sufficiently specific to hybridize to complementary template sequence during the generation of labeled nucleic acids under conditions sufficient for first strand cDNA synthesis, which conditions are known by those of skill in the art. The number of mismatches between the gene specific primer sequences and their complementary template sequences to which they hybridize during the generation of labeled nucleic acids in the subject methods will generally not exceed 20 %, usually will not exceed 10 % and more usually will not exceed 5 %, as determined using the FASTA program using default settings.

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DESCRIPTION OF THE SPECIFIC EMBODIMENTS

Arrays of polynucleotide spots and methods for their preparation are provided. In the subject arrays, a plurality of polynucleotide spots is stably associated with the surface of a solid support, where at least a portion of the polynucleotide spots on the array are made up of unique polynucleotides and all of the unique polynucleotides of the array correspond to one particular type of gene, e.g. tightly regulated human genes, genes associated with a particular disease state, genes associated with cell cycle regulation, etc. The subject arrays find particular use in gene expression assays. Also provided are sets of a representational number of gene specific primers useful in generating target nucleic acids for use with the subject arrays. In further describing the subject invention, the arrays first will be described in general terms. Next, methods for their preparation are described. Following this, a description of representative specific array types falling within the scope of the invention will be provided. Finally, a review of representative applications in which the subject arrays may be employed will be provided, where this review includes a description of the sets of a representational number of gene specific primers according to the subject invention.

Before the subject invention is further described, it is to be understood that the invention is not limited to the particular embodiments of the invention described below, as variations of the particular embodiments may be made and still fall within the scope of the appended claims. It is also to be understood that the terminology employed is for the purpose of describing particular embodiments, and is not intended to be limiting. Instead, the scope of the present invention will be established by the appended claims.

In this specification and the appended claims, the singular forms "a," "an," and "the" include plural reference unless the context clearly dictates otherwise. Unless defined otherwise, all technical and scientific terms used herein have the same meaning as commonly understood to one of ordinary skill in the art to which this invention belongs.

ARRAYS OF THE SUBJECT INVENTION-GENERAL DESCRIPTION

Array Structure

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The arrays of the subject invention have a plurality of polynucleotide spots stably associated with a surface of a solid support. Each spot on the array comprises a polynucleotide sample, i.e. polynucleotide probe composition, of known identity, usually of known sequence, as described in greater detail below. The polynucleotide spots on the array may be any convenient shape, but will typically be circular, elliptoid, oval or some other analogously curved shape. The density of the spots on the solid surface is at least about 5/cm² and usually at least about 10/cm² but does not exceed about 1000/cm², and usually does not exceed about 300/cm². The spots may be arranged in any convenient pattern across or over the surface of the array, such as in rows and columns so as to form a grid, in a circular pattern, and the like, where generally the pattern of spots will be present in the form of a grid across the surface of the solid support. See Fig. 1.

In the subject arrays, the spots of the pattern are stably associated with the surface of a solid support, where the support may be a flexible or rigid solid support. By stably associated is meant that the polynucleotides of the spots maintain their position relative to the solid support under hybridization and washing conditions. As such, the polynucleotide members which make up the spots can be non-covalently or covalently stably associated

with the support surface. Examples of non-covalent association include non-specific adsorption, binding based on electrostatic (e.g. ion, ion pair interactions), hydrophobic interactions, hydrogen bonding interactions, specific binding through a specific binding pair member covalently attached to the support surface, and the like. Examples of covalent binding include covalent bonds formed between the spot polynucleotides and a functional group present on the surface of the rigid support, e.g. -OH, where the functional group may be naturally occurring or present as a member of an introduced linking group, as described in greater detail below.

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As mentioned above, the array is present on either a flexible or rigid substrate. By flexible is meant that the support is capable of being bent, folded or similarly manipulated without breakage. Examples of solid materials which are flexible solid supports with respect to the present invention include membranes, e.g. nylon, flexible plastic films, and the like. By rigid is meant that the support is solid and does not readily bend, i.e. the support is not flexible. As such, the rigid substrates of the subject arrays are sufficient to provide physical support and structure to the polymeric targets present thereon under the assay conditions in which the array is employed, particularly under high throughput handling conditions. Furthermore, when the rigid supports of the subject invention are bent, they are prone to breakage.

The solid supports upon which the subject patterns of spots are presented in the subject arrays may take a variety of configurations ranging from simple to complex, depending on the intended use of the array. Thus, the substrate could have an overall slide or plate configuration, such as a rectangular or disc configuration. In many embodiments, the substrate will have a rectangular cross-sectional shape, having a length of from about 10 mm to 200 mm, usually from about 40 to 150 mm and more usually from about 75 to 125 mm and a width of from about 10 mm to 200 mm, usually from about 20 mm to 120 mm and more usually from about 0.1 mm to 5.0 mm, usually from about 0.1 mm to 5.0 mm, usually from about 0.1 mm to 2 mm and more usually from about 0.2 to 1 mm.

The substrates of the subject arrays may be fabricated from a variety of materials. The materials from which the substrate is fabricated should ideally exhibit a low level of non-specific binding during hybridization events. In many situations, it will also be preferable to employ a material that is transparent to visible and/or UV light. For flexible substrates, materials of interest include: nylon, both modified and unmodified, nitrocellulose,

polypropylene, and the like, where a nylon membrane, as well as derivatives thereof, is of particular interest in this embodiment. For rigid substrates, specific materials of interest include: glass; plastics, e.g. polytetrafluoroethylene, polypropylene, polystyrene, polycarbonate, and blends thereof, and the like; metals, e.g. gold, platinum, and the like; etc.

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The substrates of the subject arrays comprise at least one surface on which the pattern of spots is present, where the surface may be smooth or substantially planar, or have irregularities, such as depressions or elevations. The surface on which the pattern of spots is present may be modified with one or more different layers of compounds that serve to modify the properties of the surface in a desirable manner. Such modification layers, when present, will generally range in thickness from a monomolecular thickness to about 1 mm, usually from a monomolecular thickness to about 0.01 mm and more usually from a monomolecular thickness to about 0.001 mm. Modification layers of interest include: inorganic and organic layers such as metals, metal oxides, polymers, small organic molecules and the like. Polymeric layers of interest include layers of: peptides, proteins, polynucleic acids or mimetics thereof, e.g. peptide nucleic acids and the like; polysaccharides, phospholipids, polyurethanes, polyesters, polycarbonates, polyureas, polyamides, polyethyleneamines, polyarylene sulfides, polysiloxanes, polyimides, polyacetates, and the like, where the polymers may be hetero- or homopolymeric, and may or may not have separate functional moieties attached thereto, e.g. conjugated.

The total number of spots on the substrate will vary depending on the number of different polynucleotide probes one wishes to display on the surface, as well as the number of control spots, calibrating spots and the like, as may be desired depending on the particular application in which the subject arrays are to be employed. Generally, the pattern present on the surface of the array will comprise at least about 10 distinct spots, usually at least about 20 distinct spots, and more usually at least about 50 distinct spots, where the number of spots may be as high as 10,000 or higher, but will usually not exceed about 5,000 distinct spots, and more usually will not exceed about 3,000 distinct spots. In many embodiments, it is preferable to have each distinct probe composition presented in duplicate, i.e. so that there are two spots for each distinct polynucleotide probe composition of the array. In certain embodiments, the number of spots will range from about 200 to 600.

The amount of polynucleotide present in each spot will be sufficient to provide for adequate hybridization and detection of target nucleic acid during the assay in which the

array is employed. Generally, the amount of polynucleotide in each spot will be at least about 0.1 ng, usually at least about 0.5 ng and more usually at least about 1 ng, where the amount may be as high as 1000 ng or higher, but will usually not exceed about 20 ng and more usually will not exceed about 10 ng. The copy number of each polynucleotide in a spot will be sufficient to provide enough hybridization sites for target molecule to yield a detectable signal, and will generally range from about 0.01 fmol to 50 fmol, usually from about 0.05 fmol to 20 fmol and more usually from about 0.1 fmol to 5 fmol. Where the spot has an overall circular dimension, the diameter of the spot will generally range from about 10 to 5,000 μ m, usually from about 20 to 2,000 μ m and more usually from about 50 to 1000 μ m.

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A critical feature of the subject arrays is that at least a portion, usually the majority, of the polynucleotide spots on the array are made up of polynucleotide probes that all correspond to the same kind or kind of gene, i.e. genes that all share some common characteristic or can be grouped together based on some common feature, such as species of origin, tissue or cell of origin, functional role, disease association, etc. Other spots which may be present in the pattern include spots comprising genomic DNA, housekeeping genes, negative and positive control genes, and the like. These latter types of spots comprise polynucleotides that are not "unique" as that term is defined and used herein, i.e. they are "common." In other words, they are calibrating or control genes whose function is not to tell whether a particular "key" gene of interest is expressed, but rather to provide other useful information, such as background or basal level of expression, and the like. The percentage of spots which are made of unique polynucleotides that correspond to the same type of gene is generally at least about 30 number %, and usually at least about 60 number % and more usually at least about 80 number %. Therefore, the arrays of the subject invention will be of a specific array type, where representative array types include: human arrays, mouse arrays, cancer arrays, apoptosis arrays, human stress arrays, oncogene and tumor suppressor arrays, cell-cell interaction arrays, cytokine and cytokine receptor arrays, rat arrays, blood arrays, mouse stress arrays, neuroarrays, and the like, where some of these representative arrays are described in greater detail below.

With respect to the polynucleotide probes that correspond to a particular type or kind of gene, type or kind can refer to a plurality of different characterizing features, where such features include: species specific genes, where specific species of interest include eukaryotic

species, such as mice, rats, rabbits, pigs, primates, humans, etc.; function specific genes, where such genes include oncogenes, apoptosis genes, cytokines, receptors, protein kinases, etc.; genes specific for or involved in a particular biological process, such as apoptosis, differentiation, cell cycle regulation, cancer, aging, proliferation, etc.; location specific genes, where locations include organ, such as heart, liver, prostate, lung etc., tissue, such as nerve, muscle, connective, etc., cellular, such as axonal, lymphocytic, etc, or subcellular locations, e.g. nucleus, endoplasmic reticulum, Golgi complex, endosome, lyosome, peroxisome, mitochondria, cytoplasm, cytoskeleton, plasma membrane, extracellular space; specific genes that change expression level over time, e.g. genes that are expressed at different levels during the progression of a disease condition, such as prostate genes which are induced or repressed during the progression of prostate cancer.

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The average length of the polynucleotides on the array is chosen to be of sufficient length to provide a strong and reproducible signal, as well as tight and robust hybridization. As such, the average length of the polynucleotides of the array will typically range from about 120 to 1000 nt and usually from about 120 to 800 nt, where in many embodiments, the average length ranges from about 200 to 700 nt, and usually 200 to 600 nt. The length of each polynucleotide on the array is less than the length of the mRNA to which it corresponds. As such, the polynucleotide represents only a fraction of the full length cDNA to which it corresponds.

As mentioned above, the subject arrays typically comprise one or more additional spots of polynucleotides which do not correspond to the array type, i.e. the type or kind of gene represented on the array. In other words, the array may comprise one or more spots that are made of non "unique" polynucleotides, i.e common polynucleotides. For example, spots comprising genomic DNA may be provided in the array, where such spots may serve as orientation marks. Spots comprising plasmid and bacteriophage genes, genes from the same or another species which are not expressed and do not cross hybridize with the cDNA target, and the like, may be present and serve as negative controls. In addition, spots comprising housekeeping genes and other control genes from the same or another species may be present, which spots serve in the normalization of mRNA abundance and standardization of hybridization signal intensity in the sample assayed with the array.

Polynucleotide Probes of the Arrays

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Each spot of the pattern present on the surface of the substrate is made up of a unique polynucleotide probe composition. By "polynucleotide probe composition" is meant a collection or population of single stranded polynucleotides capable of participating in a hybridization event under appropriate hybridization conditions, where each of the individual polynucleotides may be the same -- have the same nucleotide sequence-- or different sequences, for example the probe composition may consist of 2 different single stranded polynucleotides that are complementary to each other (i.e. the two different polynucleotides in the spot are complementary but physically separated so as to be single stranded, i.e. not hybridized to each other). In many embodiments, the probe compositions will comprise two complementary, single stranded polynucleotides.

In those polynucleotide probe compositions having unique polynucleotides, the sequence of the polynucleotides are chosen in view of the type and the intended use of the array on which they are present. The unique polynucleotides are chosen so that each distinct unique polynucleotide does not cross-hybridize with any other distinct unique polynucleotide on the array, i.e. the polynucleotide of any other polynucleotide probe composition that corresponds to a different gene falling within the broad category or type of genes represented on the array. As such, the nucleotide sequence of each unique polynucleotide of a probe composition will have less than 90% homology, usually less than 85 % homology, and more usually less than 80% homology with any other different polynucleotide of a probe composition of the array, where homology is determined by sequence analysis comparison using the FASTA program using default settings. The sequence of unique polynucleotides in the probe compositions are not conserved sequences found in a number of different genes (at least two), where a conserved sequence is defined as a stretch of from about 40 to 200 nucleotides which have at least about 90% sequence identity, where sequence identity is measured as above. The polynucleotide will generally be a deoxyribonucleic acid having a length of from about 120 to 1000, usually from 120 to 700 nt, and more usually 200 to 600 nt. The polynucleotide will not cross-hybridize with any other polynucleotide on the array under standard hybridization conditions. Again, the length of the polynucleotide will be shorter than the mRNA to which it corresponds.

Array Preparation

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The subject arrays can be prepared using any convenient means. One means of preparing the subject arrays is to first synthesize the polynucleotides for each spot and then deposit the polynucleotides as a spot on the support surface. The polynucleotides may be prepared using any convenient methodology, such as automated solid phase synthesis protocols, preparative PCR and like, where preparative PCR or enzymatic synthesis is preferred in view of the length and the large number of polynucleotides that must be generated for each array.

For preparative PCR, primers flanking either side of the portion of the gene of interest will be employed to produce amplified copy numbers of the portion of interest. Methods of performing preparative PCR are well known in the art, as summarized in PCR, Essential Techniques (Ed. J.F. Burke, John Wiley & Sons)(1996). Alternatively, if a gene fragment of interest is cloned into a vector, vector primers can be used to amplify the gene fragment of interest to produce the polynucleotide.

In determining the portion of the gene to be amplified and subsequently placed on the array, regions of the gene having a sequence unique to that gene should preferably be amplified. Different methods may be employed to choose the specific region of the gene to be amplified. Thus, one can use a random approach based on availability of a gene of interest. However, instead of using a random approach which is based on availability of a gene of interest, a rational design approach may also be employed to choose the optimal sequence for the hybridization array. Preferably, the region of the gene that is selected and amplified is chosen based on the following criteria. First, the sequence that is chosen should yield a polynucleotide that does not cross-hybridize with any other polynucleotide that is present on the array. Second, the sequence should be chosen such that the polynucleotide has a low probability of cross-hybridizing with a polynucleotide having a nucleotide sequence found in any other gene, whether or not the gene is to be represented on the array from the same species of origin, e.g. for a human array, the sequence will not be homologous to any other human genes. As such, sequences that are avoided include those found in: highly expressed gene products, structural RNAs, repeated sequences found in the sample to be tested with the array and sequences found in vectors. A further consideration is to select sequences which provide for minimal or no secondary structure, structure which allows for

optimal hybridization but low non-specific binding, equal or similar thermal stabilities, and optimal hybridization characteristics.

The prepared polynucleotides may be spotted on the support using any convenient methodology, including manual techniques, e.g. by micro pipette, ink jet, pins, etc., and automated protocols. Of particular interest is the use of an automated spotting device, such as the Beckman Biomek 2000 (Beckman Instruments). As mentioned above, the polynucleotide probe compositions that are spotted onto the array surface are made up of single stranded polynucleotides, where all the polynucleotides may be identical to each other or a population of complementary polynucleotides may be present in each spot.

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SPECIFIC ARRAY TYPES OF THE SUBJECT INVENTION

A variety of specific array types are also provided by the subject invention. As discussed above, array type refers to the nature of the polynucleotide probes present on the array and the types of genes to which the probes correspond. These array types include: human array; mouse array; cancer array, apoptosis array, human stress array, oncogene and tumor suppressor arrray, cell-cell interaction array, and cytokine and cytokine receptor array, as well as other types of arrays, e.g. rat array, rat stress array, blood array, mouse stress array, and nueroarray. Each of these arrays is described separately below.

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Human Array

One specific array type provided by the subject invention is the human array. In the human array of the subject invention, the majority of the spots on the array have a polynucleotide sequence corresponding to a human gene of interest. As such, all of the unique polynucleotide probes on the array correspond to human genes. The human genes represented on the human array are typically those genes that have been identified by those of skill in the art as key genes. By "key" is meant that the genes are relevant and related to the purpose of the array, e.g. the identification of difference in the expression profiles of different cell or tissue types, where the key genes are generally functionally important to the cell. In many embodiments, the genes represented on the human array are tightly regulated human genes. The term "tightly regulated gene" is used herein in accordance with its art accepted definition and use. As such, by tightly regulated human gene is meant a gene which

is not "leaky," as opposed to housekeeping genes which are generally expressed at similar levels in different cells and different tissues, i.e. a gene which is inducible such that in response to a specific inducing signal the gene turns "on" and when this signal is removed, the gene turns "off."

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In certain embodiments of the human array, human genes that may be represented on the subject arrays include: (a) oncogenes & tumor suppressors; (b) cell cycle regulators; (c) stress response proteins; (d) ion channel & transport proteins; (e) intracellular signal transduction modulators and effectors; (f) apoptosis-related proteins; (g) DNA synthesis, repair and recombination proteins; (h) transcription factors & general DNA binding proteins; (i) growth factor & chemokine receptors; (j) interleukin & interferon receptors; (k) hormone receptors; (l) neurotransmitter receptors; (m) cell surface antigens & cell adhesion proteins; (n) growth factors, cytokines and chemokines; (o) interleukins & interferons; (p) hormones; (q) extracellular matrix proteins; (r) cytoskeleton & motility proteins; (s) RNA processing & turnover proteins; (t) post-translational modification, trafficking & targeting proteins; (u) protein turnover; and (v) metabolic pathway proteins.

In view of the length of the polynucleotides of the probe compositions of the spots, each polynucleotide of a probe composition typically has a nucleotide sequence of only a portion of the human gene. Specific sequences to which the polynucleotide sequence may correspond include those identified in Table 1 below, where by "correspond" is meant that the polynucleotide could have the same sequence as specified or a sequence complementary to the specified sequence. Whether the polynucleotide sequence is the same as a portion of the sense strand of the gene to which is corresponds or complementary thereto is based primarily on the nature of the target which the array is to be used, e.g. if the target is first strand cDNA, the polynucleotide will have a sequence found in the anti-sense DNA strand of the gene to which it corresponds.

Of particular interest is a human array of the subject invention as shown in Fig. 1. In the array, each spot on the array comprises a known polynucleotide, as specified in Table 1, where the array comprises spots which: (a) correspond to 588 different tightly regulated human genes; (b) comprise plasmid and bacteriophage polynucleotides; (c) comprise polynucleotides corresponding to housekeeping genes; and (d) genomic DNA. Each of the different types of polynucleotide spots are positioned at a known location on the membrane surface.

TABLE I

Array Coordinate	GeneBank #	Gene Name	Position
F2		interleukin-7 receptor (IL-7)	1410-1625
F.51	X01992. M29383	HUIFN-gamma interferon	391-586
FSi	. 1	interleukin 7 (IL-7)	174-447
Ala	V00568	c-myc oncogene	1372-1594
E2m	X01057, X01058, X01402	interleukin-2 receptor	1990-2247
F5k	A14844	interleukin-2 (IL-2)	181-436
E1a	M29366	epidermal growth factor receptor (ERBB3)	3886-4139
Cla	X04434, M24599	insulin-like growth factor I receptor	3414-3904
F1a	M29645	insulin-like growth factor II	436-618
Clb	L09210	homo sapiens inducible nitric oxide synthase	3503-3856
E4f	M64752	glutamate receptor subunit (GLUH1)	2232-2567
A1b	X03663	c-fms proto-oncogene	2568-2880
C1c	M32315	tumor necrosis factor receptor	3359-3543
C1d	Z12020	p53-associated gene	920-1232
F1b	X02811	platelet-derived growth factor B chain	1663-2125
B1d	X01060	transferrin receptor	4382-4770
FSI	X02851	interleukin-1 precursor (PRE IL-1)	1107-1473
F5m	K02770	monocyte interleukin 1 (IL-1)	917-1208
F5n	M14743	interleukin 3 (IL-3)	390-608
F6a	M13982	interleukin 4 (IL-4)	216-459
F6b	X04602	interleukin BSF-2 (B-cell differentiation factor)	130-555
Cle	X01394	tumor necrosis factor	607-879
C1f	D12614	lymphotoxin (TNF-BETA)	305-499
E5c	M12807	T-cell surface glycoprotein T4	947-1140
Ezn	M20566, X12830	interleukin 6 receptor	2359-2823
F6c	X04688	T-cell replacing factor (interleukin-5)	35-279
F6d	M28622	interferon beta-1 (IFN-beta-1)	345-730
F1c	M11220	granulocyte-macrophage colony stimulating factor	121-621
F1d	K03222	transforming growth factor-alpha	338-595
F6e	100209	leukocyte interferon (IFN-alpha) alpha-C	89-430
F1e	X02812, J05114	transforming growth factor-beta (TGF-beta)	2398-2575
F1f	X03438	granulocyte colony-stimulating factor (G-CSF)	901-1232
D1a	M58603	nuclear factor kappa-B DNA binding subunit	2544-3019
A1c	M15024	nucleotide sequence of the c-myb cDNA clone lambda-LMC8	1981-2176
C1a	M14694	p53 cellular tumor antigen	690-964
F1g	M19154, M22045, M22046	transforming growth factor beta-2	1538-1878

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	Position
FTh	X04571	kidney epidermal growth factor (EGF) precursor	4164-4434
E3a	J03171	interferon alpha receptor (HUIFN-ALPHA-REC)	2562-2740
F6f	M57627	interleukin 10 (IL10)	442-648
E3b	M26062	interleukin 2 receptor beta chain (P70-75)	3399-3748
E3c	M74782	interleukin 3 receptor (HIL-3RA)	651-1116
E3d	X52425	interleukin 4 receptor	2641-2974
E3e	M75914	interleukin 5 receptor alpha	555-959
E3f	X77722	interferon alpha/beta receptor	553-1012
ī	HG1621	cytokine humig	2021-2246
E4g	HG1160, M37981	cholinergic receptor nicotinic alfa polipeptide 3	934-1250
E3g	HG1252, D11086	interleukin 2 receptor gamma polipeptide	674-1006
E4b	HG1334, M20132, J03180	androgen receptor	1879-2146
E1b	HG135, M73238	ciliary neurotropic factor receptor	610-849
C1h	HG1410, X68486	adenosine receptor	1281-1494
E3h	HG1757, J03143	interferon gamma receptor	610-824
E1c	HG2246, M60459	erythropoietin receptor	1423-1740
CJ.		A1 adenosine receptor-adenylate cyclase inhibitor	508-921
B1e	- 1	orphan hormone nuclear receptor	817-1147
Ci	HG3381, X76981	adenosine receptor A3	1043-1452
E4c	L00587	calcitonin receptor	885-1270
B1f	HG74, M62424	coagulating factor II receptor	2297-2697
A1e	HG886, L07594	transforming growth factor beta receptor III 300 kDa	3358-3592
E3i	HG216, M84747	interleukin 9 receptor	289-528
E3j	HG4080, U00672	interleukin 10 receptor	2448-2803
E1d	HG423, M14764	nerve growth factor receptor	2762-3242
ESd	HG1023	Vitronectin receptor alpha subunit	2442-2473
D1b	HG125	GATA-binding protein 2	1126-1363
D1c	HG1377	CCAAT-box DNA-binding protein Hap2 homolog	958-1272
ÇİK	•	retinoic acid receptor epsilon	1315-1633
A1f	HG1470, X13293	B-myb	1873-2272
B 1g	HG1551	tyrosine kinase receptor tie	3114-3536
<u>01</u>	HG1601	tyrosine kinase receptor FLT4 class III	4236-4402
D1d	HG1603	helix-loop-helix protein 1R21	858-560
F1j	HG1650	thrombomodulin	1262-1605
D1e	HG1697	basic transcription element-binding protein 2	572-976
D1f	HG1963	basic transcription factor 62 kDa subunit	1449-1831

TABLE I (CONT)

Array Coordinate GeneBank # D1g HG 1972 E4d HG 2094 B1h HG 209 D1h HG 2158 D1i HG 244 F1k HG 2480 B1j HG 2490 B1j HG 272 D1j HG 272 B1k HG 286 B1k HG 3334 B1k HG 330 B1m HG 346 B1m HG 346 B1k HG 346 B1k HG 346 B1k HG 3509 B1k HG 3548	Gene Name helix-loop-helix protein Id-2 angiotensin II type 1a receptor alt splice 1 tyrosine kinase receptor HEK DNA-binding protein SMBP2 global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	11-382 1855-2030 2826-3144 1587-1911 1621-1886 349-657 3044-3302 2686-3053 1253-1475 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG1972 HG2094 HG2094 HG2158 HG244 HG2480 HG2480 HG2490 HG2490 HG2490 HG2892, X7520 HG2892, X7520 HG2892, X7520 HG3314 HG3314 HG3314 HG3314 HG3314 HG33183 HG3314 HG33183	helix-loop-helix protein Id-2 angiotensin II type 1a receptor alt splice 1 tyrosine kinase receptor HEK DNA-binding protein SMBP2 global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor	111-382 1855-2030 2826-3144 1587-1911 1621-1886 349-657 3044-3302 2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2094 HG209 HG2158 HG244 HG2480 HG2480 HG2480 HG2480 HG2480 HG2722 HG2811 HG2892, X7520 HG3183 HG3183 HG3183 HG3183 HG3183 HG3183 HG3183 HG348, A0978 HG346, A0978 HG3509 HG3509 HG3509 HG3509	angiotensin II type 1a receptor alt splice 1 tyrosine kinase receptor HEK DNA-binding protein SMBP2 global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor	1855-2030 2826-3144 1587-1911 1621-1886 349-657 3044-3302 2686-3053 1253-1475 1253-1475 1686-2063 2551-2820 2551-2820 2551-2820 1686-2063
HG209 HG2158 HG244 HG244 HG2480 HG2480 HG2480 HG272 HG272 HG277 HG2811 HG2892, X7520 HG2892, X7520 HG3183 HG3183 HG3183 HG3187 HG3187 HG3486 HG3486	tyrosine kinase receptor HEK DNA-binding protein SMBP2 global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	2826-3144 1587-1911 1621-1886 349-657 3044-3302 2686-3053 1253-1475 1253-1475 1686-2063 2551-2820 2551-2820 2551-2820 1686-2063 1686-2063 1686-2063 1686-2063
HG2158 HG244 HG2480 HG2480 HG2480 HG2480 HG2480 HG272 HG272 HG277 HG2811 HG2892, X7520 HG3183 HG3183 HG3183 HG346, A0978 HG346, A0978 HG3463	DNA-binding protein SMBP2 global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor	1587-1911 1621-1886 349-657 3044-3302 2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG244 HG2480 HG2480 HG2480 HG2480 HG2490 HG2722 HG2722 HG2811 K HG2892, X7520 HG3183 HG3183 HG3314 HG3187 HG3183 HG348 HG348 HG3488 HG3488 HG3488 HG3488	global transcription activator FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone trilodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor	1621-1886 349-657 3044-3302 2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2480 HG2490 HG2490 HG2490 HG272 HG2811 K HG2892, X7520 HG2892, X7520 HG3183 HG3183 HG3187 HG3187 HG3187 HG3187 HG3180 HG362 HG362 HG363 HG363	FMLP-related receptor I transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone triiodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	349-657 3044-3302 2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2490 HG2722 HG277 HG277 HG2869 K HG2869 K HG2869 HG2869 K HG2869 HG314 HG314 HG314 HG314 HG318 HG187 HG1830 HG1830 HG1830 HG187 HG369	transmembrane receptor ror1 tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone triiodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	3044-3302 2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2722 HG2811 HG2811 K HG2869 K HG2869 HG2892, X7520 HG3183 HG3183 HG314 HG3183 HG1187 HG1830 HG1830 HG1830 HG1830 HG1830 HG3630 HG3846, A0978 HG3630 HG3630 HG3630	tyrosine kinase KDR receptor DNA-binding protein ICS thyroid hormone triiodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	2686-3053 1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2811 HG2811 K HG2869 K HG2892, X7520 HG3183 HG3314 HG3314 HG183 HG187 HG187 HG187 HG1830 HG346, A0978 HG346, A0978 HG3463 HG348	DNA-binding protein ICS thyroid hormone triiodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	1253-1475 1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2811 HG2869 HG2892, X7520 HG3183 HG3314 L25124 HG1187 HG1662 HG1662 HG1662 HG3609 HG346, A0978 HG3463 HG3509 HG3510 HG3510	thyroid hormone triiodothyronine receptor c-erbA ear-1 CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	1676-2100 1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2869 HG2892, X7520 HG3183 HG3314 L25124 HG1187 HG1662 HG1830 HG1830 HG346, A0978 HG3463 HG3463 HG3509 HG3509 HG3510 HG3510	CACCC-box DNA-binding protein tyrosine kinase receptor DNA-binding protein TAX	1686-2063 2551-2820 359-765 2621-2989 1818-2029
HG2892, X7520 HG3183 HG3314 L25124 HG1187 HG1662 HG1830 HG1830 HG346, A0978 HG346, A0978 HG3463 HG3509 HG3510 HG3510	tyrosine kinase receptor DNA-binding protein TAX	2551-2820 359-765 2621-2989 1818-2029
	DNA-binding protein TAX	359-765 2621-2989 1818-2029
		2621-2989 1818-2029
	tyrosine kinase receptor TKT	1818-2029
	prostaglandin E2 receptor	
	epidermal growth factor receptor	3410-3757
	platelet-activating factor receptor	1103-1398
	tyrosine phosphatase receptor eph alt splice 1	2607-3053
	DNA-binding protein/plasminogen activator inhibitor-1 regulator	1304-1736
	interferon gamma receptor	66-317
	DNA-binding protein CN sterol regulating	96-341
	v-erbA related ear-2 protein	882-1057
8	v-erbA related ear-3 protein	1449-1700
	CCAAT displacement protein cut homolog alt splice 1	2000-2400
D2b HG3748	basic transcription factor 44 kDa subunit	606-843
	DNA-binding protein APRF	1545-1575
	estrogen receptor hSNF2b	2415-2682
B2a HG4196	urokinase-type plasminogen activator receptor	749-1043
A1i HG4269	Ets-like gene	710-1064
B2b HG4279	tyrosine kinase TRK-B receptor	1006-1384
	DNA-binding protein NFX1 cysteine-rich specific	2003-2311
	DP2 dimerization partner of E2F	1603-1838
F11 HG563	glia maturation factor beta	203-434
	DNA-binding protein TAXREB67	1059-1495
D20 HG859, L05515	cAMP-responsive element-binding protein	807-1120
	tyrosine kinase EGF receptor Her4	3570-3965

TABLE I (CONT)

Arran Coordinate	ConoBank #	Gene Name	Position
Ariay coolument	_	tyrosine phosphatase receptor gamma polypeptide	3623-3938
720	HG970	DNA-binding protein PO-GA	3196-3413
120	HG99 M64673	CCAAT enhancer-binding protein beta	294-572
A11	J04111	c-jun proto-oncogene (jun) clone HCJ-1	2207-2583
E3	M27492	interleukin 1 receptor	3847-4288
C1m	M33294	tumor necrosis factor receptor	1570-1817
F1m	M37435	macrophage-specific colony-stimulating factor (CSF-1)	2277-2413
A1m	YOO285	insulin-like growth factor II receptor	1394-1831
A1n	HG404	tyrosine kinase receptor HER2	2556-2722
B2d	D10923	HM74	1357-1826
B2e	D10924	HM89	351-808
BZÍ	D10925	HM145	1353-1832
Fin	D14012	hepatocyte growth factor activator precursor	1487-1845
F2a	D16431	hepatoma-derived growth factor	359-625
F2h	D30751	bone morphogenetic protein 4	943-1321
B2a	J03358	FER tyrosine kinase	2384-2688
For	J04130	activation (Act-2)	236-592
F2d	J05081	endothelin ET3	1428-1685
F28	K03515	neuroleukin	1368-1656
Δ2a	1 06139	TEK tyrosine kinase receptor	3243-3586
F10	L06622	endothelin receptor EDNRA	870-1080
11 H	L06623	endothelin receptor EDNRB	497-814
F6a	L06801	interleukin IL-13	285-743
C10	L07414	CD40 ligand	863-1277
C2a	L08096	CD27 ligand	233-627
E3m	L08187	cytokine receptor (EB13)	627-1019
F2f	L12260	glial growth factor 2 (recombinant)	1069-1452
F20	L12261	glial growth factor (recombinant)	762-1041
FSh	L15344	interleukin IL-14	1181-1562
F2h	L36052	thrombopoietin (MGDF/Mpl ligand)	230-613
<u>F1</u>	M10051	insulin receptor	3274-3758
F2i	M17778	uromodulin	1463-1913
F2;	M21121	RANTES pro-inflammatory cytokine	180-545
15	M21574	PDGF-alpha receptor	5118-5583
E1K	M21616	PDGF-beta receptor	842-1133
101	M22488	bone morphogenetic protein 1	702-1098

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	Position
	M22489	bone morphogenetic protein 2a	267-997
F2m	M22491	bone morphogenetic protein 3	1458-1731
F2n	M23452	macrophage inflammatory protein GOS19-1	243-704
F3a	M24545	monocyte chemotactic and activating factor MCAF	36-384
F3b	M25667	neuronal growth protein GAP-43	747-1154
F3c	M27288	oncostatin M	833-1113
F3d	M30704	amphiregulin AR	511-837
F3e	M31145	insuline-like growth factor binding protein 1	476-861
E11	M31165	TNF-inducible hyaluronate-binding protein TSG-6	320-584
F3f	M32977	heparin-binding vascular endothelial growth factor VEGF	198-622
A2b	M35410	insuline-like growth factor binding protein 2	680-1071
F7a	M36717	ribonuclease/angiogenun inhibitor RAI	713-1028
F3a	M37722	bFGF receptor	1746-1967
B2h	M57230	glycoprotein gp130	1757-2152
F3h	M57399	nerve growth factor HBNF-1	602-847
F3.	M57502	secreted protein I-309	205-397
F6i	M57765	interleukin IL-11	132-460
Elm	M59818	granulocyte colony-stimulating factor receptor G-CSFR1	1453-1891
F3	M59964	stem cell factor	898-1283
F3K	M60278	heparin-binding EGF-like growth factor	1905-2146
F3I	M60718	HGF (hepatocyte growth factor)	1549-1970
F3m	M60828	keratinocyte growth factor	419-766
F3n	M61176	brain-derived neurotrophic factor BDNF	982-1265
F4a	M62302	growth/differentiation factor GDF-1	615-957
Ein	M62505	C5a anaphylatoxin receptor	725-1098
E5e	M63928	T cell activation antigen CD27	513-977
F4b	M65199	endothelin ET2	338-570
F6i	M65290	interleukin IL-12 (NKSF p40)	622-848
Fé	M65291	interleukin IL-12 (NKSF p35)	066-009
C2b	M67454	Fas antigen	2063-2288
E2a	M68932	interleukin 8 receptor alpha (IL8RA)	1179-1370
E2b	M73482	NMB-R (neuromedin B receptor)	282-544
F4c	M74178	hepatocyte growth factor-like protein	1643-2015
A5c	M76125	AXL tyrosine kinase receptor	2054-2328
E5f	M83554	lymphocyte activation antigen CD30	3152-3421
	1000004	this contract to	40-342

TABLE I (CONT)

		Nome Nome	Position
Array Coordinate	-+		4 450 4749
F4e	M92934	connective tissue growth factor	1439-1740
C2c	M93426	tyrosine phosphatase receptor zeta-polypeptide	5090-1748
F4f	M96956	TDGF3	1294-1712
E2c	S59184	RYK=related to receptor tyrosine kinase isolog	1760-1968
A2c	U01134	VEGF receptor	1288-1604
F2d	U01839	Duffy blood group antigen (Fya-b+)	127-150
ASd	U02687	growth factor receptor tyrosine kinase STK-1	2491-2965
F3n	U03187	interleukin 12 receptor component	1053-1381
		monocyte chemoattractant protein 1 receptor (MCP-1RA) alternatively	
E2e	U03882	spliced	1514-1799
		monocyte chemoattractant protein 1 receptor (MCP-1RB) alternatively	1262 1713
E2f	U03905	spliced	13051
C2d	U04806	FLT3/FLK2 ligand	29-362
F40	U10117	endothelial-monocyte activating polypeptide II	272-304
E20	U11814	keratinocyte growth factor receptor	753-1189
C2e	U13737	cysteine protease CPP32 isom alpha	2007-2434
F6I	U14407	interleukin IL-15	338-695
F2h	U14722	activin type I receptor	333-740
F4h	U43142	VRP (vascular endothelial growth factor related protein)	1165-1559
F4;	X02530	IFN-gamma-inducible chemokine IP-10	280-613
A1d	X06182	c-kit proto-oncogene	37-430
EA:	X06233	MRP-14 (calcium binding protein in macrophages MIF-related)	16-254
APH.	X06234	MRP-8 (calcium binding protein in macrophages MIF-related)	37-351
F41	X06374	platelet-derived growth factor A chain PDGF-A	522-955
F4m	X13967	leukemia inhibitory factor LIF	1810-2239
F6m	X17543	interleukin IL-9 (P40)	156-186
E2i	X17648	granulocyte-macrophage colony-stimulating factor receptor GM-CSFRa	868-1173
F4n	X51943	fibroblast growth factor FGF-1	1131-1502
F5a	X53655	nerve growth factor NGF-2 (same as NT-3)	112-416
FSh	X53799	macrophage inflammatory protein-2alpha (MIP2alpha)	157-501
F50	X54936	PIGF (placenta growth factor)	1098-1371
F43	X59770	interleukin 1 receptor type II	842-1244
F2i	X60592	Cdw40	198-605
F2X	X72304	beta-thromboglobulin-like protein	230-533
ESd	X78686	neutrophil-activating peptide ENA-78	65-329
8	6265ZX	OX40 ligand/qp34	329-657

TABLE I (CONT)

Array Coordinate	Generalik #		
F5f	_	monocyte-derived neutrophil chemotactic factor MDNCF	99-287
B2i	D10495	protein kinase C delta-type	1467-1817
D2i	D13316	transcription factor E4TF1-47	965-1175
D2k	D13318	transcription factor E4TF1-60	1069-1512
CSi	D13804	recA-like protein HsRad51	867-1159
E5q	D13866	alpha-catenin	2235-2577
A5e	D13889	H[-p]	83-433
D2i	D15050	transcription factor AREB6	2417-2680
C2f	D15057	DAD-1	124-334
A2d	D17517	sky Sky	2132-2597
B2i	D21878	BST-1	206-980
D2m	D26120	ZFM1 protein	2367-2704
D2n	D26121	ZFM1 protein alternatively spliced product	440-908
D3a	D26155	transcriptional activator hSNF2a	3917-4258
B2k	D26309	LIMK (LIM kinase)	2810-3157
D3b	D28118	DB1	1166-1481
D3c	D28468	DNA-binding protein TAXREB302	386-811
Esh	J03132	intercellular adhesion molecule-1 (ICAM-1)	1220-1599
A2e	J03241	transforming growth factor-beta 3 (TGF-beta3)	1416-1833
F7b	J03634	erythroid differentiation protein (EDF)	983-1372
ESi	J04536	sialophorin (CD43)	178-392
CSi	L04791	excision repair protein ERCC6	1772-2194
BZI	L05624	MAP kinase kinase	842-1217
CSK	L07540	replication factor C 36-kDa subunit	708-1051
CSI	L07541	replication factor C 38-kDa subunit	438-762
D3d	L08424	achaete scute homologous protein (ASH1)	1113-1455
AZÍ	L11353	moesin-ezrin-radixin-like protein	355-674
D3e	L11672	Kruppel related zinc finger protein (HTF10)	107-555
B2m	L13616	focal adhesion kinase (FAK)	2179-2631
B2n	L13738	activated p21cdc42Hs kinase (ack)	758-1184
ASÍ	L13740	TR3 orphan receptor	818-1077
D3f	L14611	transcription factor RZR-alpha	620-982
A2a	L14837	tight junction (zonula occludens) protein ZO-1 (tumor suppressor)	6327-6660
C2a	L16785	c-myc transcription factor (puf)	69-351
B3a	L19067	NF-kappa-B transcription factor p65 subunit	1897-2137
0.74	1 10185	natural Villar cell enhancing (actor (NKEFR)	348-736

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	Position
D3a		paired box homeotic protein (PAX8)	113-338
CSm	L20046	ERCC5 excision repair protein	1374-1638
B3b	L20320	protein serine/threonine kinase stk1	89-305
B3c	L20321	protein serine/threonine kinase stk2	2534-2802
B3d	L20422	14-3-3n protein	163-671
D3h	L20433	octamer binding transcription factor 1 (OTF1)	3275-3583
ESi	L20815	S protein	1677-2107
Bla	L20977	plasma membrane calcium ATPase isoform 2 (ATP2B2)	3861-4236
B3e	L22075	guanine nucleotide regulatory protein (G13)	1073-1376
C2h	L22474	Bax beta	227-278
CSn	L24564	Rad	489-780
B3f	L24959	calcium/calmodulin dependent protein kinase	969-1220
B30	L25259	CTLA4 counter-receptor (B7-2)	496-722
ig	L29511	GRB2 isoform	355-573
D3i	L31881	Inuclear factor I-X	415-729
B3h	L32976	protein kinase (MLK-3)	970-1283
ASa	L33264	CDC2-related protein kinase (PISSLRE)	454-755
D3i	L34587	RNA polymerase II elongation factor SIII p15 subunit	115-354
B3i	L35233	autocrine motility factor receptor (AMFR)	1221-1514
Azh	M13150	mas proto-oncogene	262-726
D3k	M14631	guanine nucleotide-binding protein G-s alpha subunit partial cds	824-1120
B1b	M15800	MAL protein	461-695
Dai	M16937	homeobox c1 protein	367-667
ESK	M21097	differentiation antigen (CD19)	740-1071
B3i	M22199	protein kinase C alpha-polypeptide (PKCA)	767-1106
ESI	M23197	differentiation antigen (CD33)	885-1141
ASh	M26708	prothymosin alpha (ProT-alpha)	538-864
B3k	M28210	GTP-binding protein (RAB3A)	288-591
83	M28211	GTP-binding protein (RAB4)	255-495
ВЗт	M28212	GTP-binding protein (RAB6)	59-310
B3n	M28213	GTP-binding protein (RAB2)	56-269
B4a	M28214	GTP-binding protein (RAB3B)	322-621
B4b	M28215	GTP-binding protein (RAB5)	447-672
A5i	M28882	MUC18 glycoprotein	1756-2180
D3m	M29038	stem cell protein (SCL)	2804-3086
100	M29142	myeloblastin	312-693

TABLE ! (CONT)

Array Coordinate	Cana Rank #	Gene Name	Losition
F5m	M30257	vascular cell adhesion molecule 1	1056-1450
E5n	M30640	endothelial leucocyte adhesion molecule I (ELAM1)	2098-2549
C6a	M30938	Ku (p70/p80) subunit	2340-2764
A2i	M31213	papillary thyroid carcinoma-encoded protein	2285-2631
D3n	M31523	transcription factor (E2A)	2277-2685
B4c	M31630	cyclic AMP response element-binding protein (HB16) 3' end	316-636
C6b	M31899	DNA repair helicase (ERCC3)	2109-2466
Cec	M32865	Ku protein subunit	1729-1974
E6a	M33374	cell adhesion protein (SQM1)	53-354
E6b	M34064	N-cadherin	942-1299
B4d	M34356	active transcription factor CREB	433-780
D4a	M34960	transcription factor IID	561-843
C6d	M36089	DNA-repair protein (XRCC1)	1226-1539
B4e	M36429	transducin beta-2 subunit	443-789
B4f	M36430	transducin beta-1 subunit 3' end	58-338
D4b	M36542	lymphoid-specific transcription factor	647-942
D4c	M36711	sequence-specific DNA-binding protein (AP-2)	950-1211
A2i	M54915	h-pim-1 protein (h-pim-1)	893-1187
E6c	M54992	B cell differentiation antigen	963-1224
E6d	M59040	cell adhesion molecule (CD44)	1158-1408
A2k	M60915	neurofibromatosis protein type I (NF1)	740-1027
D4d	M62397	colorectal mutant cancer protein	3626-3902
D4e	M62810	mitochondrial transcription factor 1	640-668
D4f	M62829	transcription factor ETR103	989-1276
D40	M62831	transcription factor ETR101	1018-1410
Cée	M63488	replication protein A 70kDa subunit	1498-1838
A5k	M63618	bullous pemphigoid antigen	5680-6055
D4h	M63896	transcriptional enhancer factor (TEF1) DNA	2935-3238
E6e	M74387	cell adhesion molecule L1 (L1CAM)	3197-3483
Cef	M74524	HHR6A (yeast RAD 6 homologue)	175-433
E6f	M74777	dipeptidyl peptidase IV (CD26)	1205-1507
CZi	M74816	sulfated glycoprotein-2 3'end	709-990
D4i	M75952	homeobox protein (HOX-11)	1209-1552
D4i	M76541	DNA-binding protein (NF-E1)	706-1053
D4K	M76766	transcription factor (TFIIB)	407-769
	100003	UEB holiv loop holiv protoin (HEB)	2676, 2084

TABLE I (CONT)

ı			Donition
Array Coordinate	GeneBank #	Gene Name	POSITION POS
D4m	M81601	transcription elongation factor (SiI)	227-293
42	M81750	myeloid cell nuclear differentiation antigen	549-873
ASI	M81757	S19 ribosomal protein	113-408
D4n	M81840	NRL gene product	946-1158
D5a	M83234	nuclease-sensitive element DNA-binding protein	790-1099
200	M84820	retinoid X receptor beta (RXR-beta)	643-1135
Cen	M87338	replication factor C 40-kDa subunit (A1)	882-1286
60 C	M87339	replication factor C 37-kDa subunit	98-355
190 190 190 190 190 190 190 190 190 190	M87503	FN-responsive transcription factor subunit	1057-1520
D5c	M92299	homeobox 21 protein (HOX2A)	1718-1945
DSd	M92843	zinc finger transcriptional regulator	892-1271
D.50	M93255		728-1118
FA9	M95489	follicle stimulating hormone receptor	1507-1752
Daf	M96824	nucleobindin precursor	701-1068
250	M96944	B-cell specific transcription factor (BSAP)	2446-2771
25.0	M97287	MAR/SAR DNA binding protein (SATB1)	1921-2226
75. 75.	M97676	(region 7) homeobox protein (HOX7)	1091-1450
FAh	S64045	5HT1a=5-hydroxytryptamine receptor (transmembrane regions 5 and 6)	128-413
A5m	101160	transmembrane 4 superfamily protein (SAS)	98-409
BAG	102081	quanine nucleotide regulatory protein (NET1)	1079-1323
RAh	102082	quanine nucleotide regulatory protein (tim1)	1852-2185
051	1102326	clone ndf43 neu differentiation factor	1430-1701
0.51	1102368	PAX3/forkhead transcription factor fusion	2231-2569
251	U02619	TFIIIC Box B-binding subunit	5023-5369
D5m	U02683	alpha palindromic binding protein	1630-2062
A2m	U03056	tumor suppressor (LUCA-1)	2039-2444
DSn	U03494	transcription factor LSF	1358-1681
B4i	U03688	dioxin-inducible cytochrome P450 (CYP1B1)	1212-1556
D6a	U04847	Init	125-538
Deb	U05040	FUSE binding protein	1002-1339
A50	U05340	p55CDC	1236-1522
84	U05875	clone pSK1 interferon gamma receptor accessory factor-1 (AF-1)	1702-2039
B1c	U07139	voltage-gated calcium channel beta subunit	2008-2383
B4k	U07236	mutant lymphocyte-specific protein tyrosine kinase (LCK)	930-1207
A6a	U07616	amphiphysin	1740-2143
i Pa	1007207	epidermal growth factor receptor substrate (eps15)	1828-2140

TABLE I (CONT)

		N. T.	0001100
Array Coordinate	GeneBank #	Gene Name	Position
E6g	U07819	contactin 1 precursor (CNTN1)	2735-3130
Dec	U08015	NF-ATc	2039-2374
P9Q	U08191	R kappa B	4657-4920
Dee	U08853	transcription factor LCR-F1	1575-1928
B4m	U09564	serine kinase	487-833
Def	009579	melanoma differentiation associated (mda-6)	1745-2063
B4n	109607	JAK family protein tyrosine kinase JAK3	3556-3892
Deg	U10323	nuclear factor NF45	967-1380
Deh	U10324	nuclear factor NF90	2901-3146
D6i	U10421	HOX A1 homeodomain protein (HOXA1)	132-492
Dei	U12535	epidermal growth factor receptor kinase substrate (Eps8)	2293-2645
C2I	U13021	positive regulator of programmed cell death ICH-1L (Ich-1)	851-1218
D6k	U13897	homolog of Drosophila discs large protein isoform 1 (hdlg-1)	2248-2624
Dei	U14575	(ard-1)	665-942
D6m	U14755	LIM domain transcription factor LIM-1 (hLIM-1)	479-759
Den	U15979	(dlk)	1090-1403
B5a	U16031	transcription factor IL-4 stat	1816-2118
Cei	X06745	DNA polymerase alpha-subunit	3721-4093
A2n	X07024	X chromsome CCG1 protein inv in cell proliferation	4002-4343
A3a	X15218	ski oncogene	2354-2662
A3b	X15219	sno oncogene snoN protein ski-related	2224-2652
E6h	X16841	N-CAM (a nontransmembrane isoform) from skeletal muscle	2338-2646
A3c	X51630	Wilms tumor WT1 zinc finger protein Krueppel-like	1866-2254
D7a	X55122	GATA-3 transcription factor	1097-1383
A6b	X55504	P120 antigen	1970-2245
D7b	X59738	ZFX put transcription activator isoform 1	749-1113
D7c	X67951	proliferation-associated gene (pag)	543-856
85b	X70326	MacMarcks	638-1008
B5c	X74979	TRKE	2138-2411
E6i	Z26317	desmoglein 2	2819-3135
F7c	A00914	angiotensin-converting enzyme (ACE)	2123-2483
F7d	A06925	relaxin H2	123-427
F7e	D10232	renin-binding protein	289-589
E4i	D28538	glutamate receptor type 1 subtype 5a	3745-4027
F7f	J04040	glucagon	201-540
E4i	L19058	glutamate receptor 5	2514-2779

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	Position
	M13981	inhibin A-subunit	828-1183
F7h	M14200	diazepam binding inhibitor	67-257
F4k	M15169	Beta-2-adrenergic receptor	2412-2783
E41	M29066	dopamine d2 receptor	1226-1521
F7i	M31159	growth hormone-dependent insulin-like growth factor-binding protein	451-744
F7i	M68867	retinoic acid-binding protein II	489-863
E4m	M76446	alpha A1 adrenergic receptor	1599-1942
E4n	M86841	serotonin receptor type 2	938-1239
F7k	U06863	follistatin-related protein precursor	1093-1425
F71	X58022	corticotropin-releasing factor-binding protein	853-1140
A6c	HT0121	cyclin-dependent kinase 2	1774-2180
A6d	HT0191	cell division cycle protein 25A tyrosine phosphatase	1632-1978
A6e	HT0285	cyclin D3	537-894
Céi	HT330	single-stranded DNA-binding protein pur-alpha	563-855
A6f	HT0609	cyclin A	876-1218
Cek	HT767	DNA topoisomerase I	2388-2796
Cel	HT784	DNA topoisomerase II alpha	2459-2883
Côm	HT1104	6-O-methylguanine-DNA methyltransferase	241-546
Cen	HT1175	DNA excision repair protein ERCC2 5' end	1520-1821
A3d	HT1426	prohibitin	172-455
A3e	HT1436	proto-oncogene raf	1704-1989
CSm	HT1483	glutathione reductase	719-1057
A3f	HT1489	proto-oncogene c-abl tyrosine protein kinase alt transcript 1	3240-3612
Ago	HT1547	cyclin D1	3427-3784
CSu	HT1790	glutathione S-transferase 12	72-420
CZa	HT1848	DNA excision repair protein ERCC1 alt transcript 1	625-938
C3a	HT2041	glutathione S-transferase M1	504-906
C3b	HT2042	glutathione S-transferase pi	203-511
SS	HT2168	glutathione S-transferase A1	257-583
A6h	HT2181	cyclin D2	3932-4284
A3a	HT2291	proto-oncogene c-src1 tyrosine kinase domain	893-1189
A3h	HT2788	proto-oncogene rel	1357-1605
A3i	HT2856	proto-oncogene rhoA multidrug resistance protein	290-572
C3d	HT2859	glutathione peroxidase	454-745
A3i	HT3039	proto-oncogene shb src-2 homolog	1365-1657
	00 FOL		412,676

TABLE I (CONT)

-4	C-1000mt #	Sone Name	Position
Array Coordinate		superoxide dismutase 1 cytosolic	198-486
273	HT3337	DNA mismatch repair protein hmth1	1765-2020
0/0	11 3337	cell division cycle protein 25 nucleotide exchange factor	3372-3651
Abi	110410	tumor suppressor DCC coloractal	3749-4042
A3k	H13563	Machine DASO rediction	789-1082
C3I	H13514	veryderms nigmentosum group C repair complementing protein	
7.0	HT/200	Action in pignical coopies of coo	582-885
270	HT4247	xeroderma pigmentosum group C repair complementing protein HHR23A	355-632
200	HTASAO	Cyclin H	717-1026
207	HT4547	olutathione S-transferase T1	617-914
250	HTE168	ionizing radiation resistance-conferring protein	856-1114
เรา เรา	100200	andothelial membrane alveoprotein IIIA (GPIIIA)	2038-2373
100	104145	neutrophi adherence receptor alpha-M subunit	2888-3183
TOK	105699	integrin beta-5 subunit	2279-2528
io i	20000	integrin alpha 4 subunit	2709-3063
EQU	L 12002	leukocyte adhesion protein (LFA-1/MAC-1/P15095 family) beta subunit	2367-2664
Ebn	W (2000)	natalet alvonotein IIB (GPIB)	268-639
E7a	M34480	integrin B-6	1619-1901
E/0		integrin alpha-3 chain	2562-2944
E/c	MOSSI I	la konde adhesion alvoorotein P15095	88-271
E/0		(ibronectin recentor alpha subunit	2094-2367
E/8	X00230	fibronectin recentor heta subunit	2116-2482
E71	VU/9/9	integrin alpha 6	3642-3988
E7g	X53565	integrin dela d	5357-5697
E/n	X3330/	integrin alpha subjunit	2690-2976
E/1	V24005	alpha 7B integrin	255-591
E/ j	V00796	leukocyte-associated molecule-1 alpha subunit (LFA-1 alpha subunit)	4526-4856
E/8	D38122	Fas ligand	516-840
B7;	D49547	heat-shock protein 40	1400-1782
D74	J03133	transcription factor SP1 3' end	1876-2272
200	107032	protein kinase C theta (PKC)	2306-2601
Bro	1 26318	protein kinase (JNK1)	952-1263
200	1 97911	CDK4-inhibitor (p16-INK4)	482-836
BSf	L35253	p38 mitogen activated protein (MAP) kinase	925-1204
B50	L36719 ·	MAP kinase kinase 3 (MKK3)	790-1169
S.		MAND Lingue Lingue 4 (MKKA)	2788-3103

TABLE I (CONT)

Array Coordinate GeneBank #	GeneBank #	Gene Name	Position
C3i	M13228	N-myc oncogene protein	761-1188
A3i	M15400	retinoblastoma susceptibility	2839-3101
A3m	M15990	c-yes-1	1325-1676
BSi	M16038	"lyn, tyrosine kinase"	1393-1666
A3n	M19720	L-myc protein	5847-6118
A4a	M19722	fgr proto-oncogene encoded p55-c-fgr protein	521-856
A6I	M25753	cyclin B	979-1311
85i	M27545	protein kinase C (PKC) type beta I	1561-1821
BSK	M31158	CAMP-dependent protein kinase subunit RII-beta	1305-1506
87;	M34664	chaperonin (HSP60)	533-839
BSI	M35203	protein-tyrosine kinase (JAK1)	2768-3054
C71	M60974	growth arrest and DNA-damage-inducible protein (gadd45)	526-886
B5m	M65066	cAMP-dependent protein kinase regulatory subunit RI-beta 3' end	444-662
A6m	M73812	cyclin E	1295-1658
A4b	M74088	APC	7992-8326
D7e	M83221	I-Rel	853-1129
BSn	M84489	extracellular signal-regulated kinase 2	1241-1522
D71	M97190	Sp2 protein	396-682
D7a	M97191	Sp3 protein	1588-1987
C70	S40706	GADD153=growth arrest and DNA-damage-inducible	480-789
ř	U25994	cell death protein (RIP)	848-1123
B6a	U30473	putative src-like adapter protein (SLAP)	524-901
CZh	U35835	DNA-PK	2250-2680
A6n	U40343	CDK inhibitor p19INK4d	750-952
E71	U43522	cell adhesion kinase beta (CAKbeta)	3658-3952
A4c	U43746	breast cancer susceptibility (BRCA2)	10056-10346
A7a	U47413	cyclin G1	755-1035
A7b	U47414	cyclin G2	989-1254
A7c	U66838	cyclin A1	1205-1456
A4d	X02751	N-ras	708-1064
B7k	X07270	heat shock protein hsp86	380-577
B6b	X07767	cAMP-dependent protein kinase catalytic subunit type alpha (EC 27137)	460-740
A4e	X16706	fra-2	376-663
A4f	X16707.	fra-1	617-897
A40	X51521	ezrin	1611-1883
B71	X54079	heat shock protein HSP27	423-683

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	Position
Dec	X54637	tyk2 non-receptor protein tyrosine kinase	3787-4110
444	Y56681	Quni	508-780
A4n	VEOGS	C-src-kinase	488-876
A4!	A33332	FBK1 protein serine/threonine kinase	754-1094
Red	VOO.000	FERK3	806-1267
Bee	X80692	ENIX.	865-1239
C3I	X86779	FASI KIIdase	2061-2463
E7m	X87838	Deta-catenin	035-1200
C3m	X89986	NBK apoptotic inducer protein	20,027
A7d	X92669	p35 cyclin-like CAK1-associated protein	2024 2003
B6f	Z29090	phosphatidylinositol 3-kinase	3021-3203
C3n	L11015	lymphotoxin-beta	03-459
Bhn	L31951	protein kinase (JNK2)	030-1000
Beh	L34583	tyrosine phosphatase (clone HFAP10)	13/2-1/01
743	141690	TNF receptor-1 associated protein (TRADD)	1009-1313
245	MAAZAE	pd-2	5087-5382
25	1145470	NIP1 (NIP1)	412-719
240	010172	(NID3 (NID3)	272-637
249	1100007	Cysteine professe MCH2 isom beta (MCH2)	387-697
C4e	UZUBBY	DAV Aratain	1371-1661
C4f	U23765	DAN protein	763-1107
C4g	U28014	Cysteme protesse (ionitre ii)	64-293
C4h	U29680	A1 protein	1018-1413
B6i	U34819	JNK3 alphaz protein kinase (JINN3AZ)	1444-1848
C4i	U45878	inhibitor of apoptosis protein 1	2000-2363
C4i	U45879	inhibitor of apoptosis protein 2	266-621
C4k	U45880	X-linked inhibitor of apotosis protein XIAP	986.1289
2	U56390	cysteine protease ICE-LAPb	211-616
C4m	U57059	Apo-2 ligand	2276-2690
C4n	U60519	apoptotic cysteine protease Mcn4 (Mcn4)	1307,1607
C5a	U60520	apoptotic cysteine protease Mcho Isom alpira (Mcho)	478-695
BGi	X14454	interferon regulatory factor 1	2440.9796
CSb	X96586	FAN protein	277.702
CFC	Y09392	WSL-LR WSL-S1 and WSL-S2 proteins	1407-1671
075	011117	homeobox HOX 4A homeodomain protein	4200-444/
A70	D38305	Tob	626-926
730	D42108	phospholipase C	1635-2003
DOR 7.2	D45130	zinc-finger DNA-binding protein	5113-5551
170	201010		

TABLE I (CONT)

Array Coordinate	GeneBank #	Gene Name	
Allay Cooluliate	D40304	serotonin 5-HT3 receptor	1703-2000
E3a	1 16/16/4	ETS oncodene (PEP1)	418-711
74	1 20216	CK2	1106-1356
A/I	1,00000	C K3	551-1002
A7g	129220	23.0	144-459
A7h	129222	OLN-	2097-2395
E5b	L76224	NMUA receptor	1962-2225
87m	M11717	heat shock protein (HSP /U)	659.919
F50	M27544	insulin-like growth factor	0025-019
Bel	M68516	protein C inhibitor	9032-9423
F.5h	M86528	neurotrophin-4 (NT-4)	6/01-17/
Bem	U09578	MAPKAP kinase (3pK)	480-83/
Δ7;	U10564	CDK tyrosine 15-kinase WEE1HU (WEE1HU)	7001-6071
22	U12134	DNA damage repair and recombination protein RAD52	1528-1/33
Ren	U14187	receptor tyrosine kinase ligand LERK-3 (EPLG3)	000-0/1
B79	1114188	receptor tyrosine kinase LERK-4 (EPLG4)	168-436
470	1118087	3'5'-cAMP phosphodiesterase HPDE4A6	1119-1453
7.70	1121002	CD40 receptor associated factor 1 (CRAF1)	980-1322
230	1122398	CDK-inhibitor P57KIP2 (KIP2)	1048-1316
W/	1124166	FB1	488-796
A4K	024100	a. a.	3054-3444
A41	07000	transcription factor TFIIIB 90 kDa subunit (HTFIIIB90)	2336-2605
D7j	UZBB3B	transcription initiation factor TEIID subunit TAFII31	260-638
D7k	M30504	II 17	257-578
F6n	U32659	orthonic dinain light chain 1 (helic1)	48-265
CSe	U32944	Cytoplashiic dynam light chair ((1905)	3507-3784
B7c	U33635	COION CARCILIONIA MINASE** (CONT.)	8938-9135
C7j	U33841	ataxia telafigieciasia (ATM)	1072-1391
A7k	U35735	KACHII (NACHII)	541-844
CSf	U39613	Cysteine processe for the cystein businesse (MIKKE)	1060-1389
B7d	U39657	MAP Milase Milase O (Milase)	1245-1530
B7e	040282	ווופקווודיווואכט איוומספ (יביג)	143-356
A7I	U41816		1967-2400
D71	U43188	Ets transcription factor (NEMF-2)	1455.1849
B7f	U43408	tyrosine kinase (TRKT)	1417.1679
A4m	U57456	transforming growth factor-beta signaling protein-1 (DSP-1)	121.403
CSg	U59747	Bcl-w (bcl-w)	674-887
1	000011	HOAT into continue protein I I DAT	\000±

TABLE I (CONT)

Array Coordinate GeneBank #	GeneBank #	Gene Name	Position
E7n	U60800	semaphorin (CD100)	2517-2921
A4n	U61262	neogenin	3144-3573
C7k	U63139	Rad50 (Rad50)	5117-5435
A5a	U68162	thrombopoietin receptor (MPL)	2184-2448
CSh	U71364	serine proteinase inhibitor (P19)	618-986
C71	X83441	DNA ligase IV	2787-3074
C7m	X84740	DNA ligase III	2460-2780
C7n	X90392	DNase X	2038-2427
B7n	HT4197	glutaredoxin	43-325
F7m	U08098	estrogen sulfotransferase (STE)	533-852
F7n	X54469, M28019	beta-preprotachykinin	321-7888
B7g	L25876	protein tyrosine phosphatase (CIP2)	110-499
A7m	M81934	CDC25B	2286-2602
A7n	U17075	P14-CDK inhibitor	116-462
G12	X01677	LIVER GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE	663-932
G13	K00558	TUBULIN ALPHA	
		HLA CLASS I HISTOCOMPATIBILITY ANTIGEN, C-4 ALPHA CHAIN	
G14	M11886	[MHC]	
G19	X00351	BETA-ACTIN	692-1077
G20	X56932	23 kDa HIGHLY BASIC PROTEIN	*
G21	U14971	RIBOSOMAL PROTEIN S9	
GS	M26880	UBIQUITIN	1922-2181
95	M86400	PHOSPHOLIPASE A2	
25	V00530	HYPOXANTHINE-GUANINE PHOSPHORIBOSYLTRANSFERASE	

Mouse Array

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In the mouse array according to the subject invention, all of the unique polynucleotide probe compositions will correspond to a mouse gene of interest. Mouse genes that are represented on the array are key genes, by which is meant that they have been reported to play primary roles in a variety of different biological processes. Typically the mouse genes represented on the array are genes that are under tight transcriptional control. Genes of interest that may be represented on the array include: oncogenes, cell cycle genes, apoptosis genes, growth factor genes, cytokine genes, interleukin genes, receptor genes, and genes associated with different stages of embryonic development.

In certain embodiments, of particular interest is an array having the following types of genes represented on its surface: oncogenes & tumor suppressors; cell cycle regulators; stress response proteins; ion channel & transport proteins; intracellular signal transduction modulators & effectors; apoptosis-related proteins; DNA synthesis, repair & recombination proteins; transcription factors & general DNA binding proteins; growth factor & chemokine receptors; interleukin & interferon receptors, hormone receptors; neurotransmitter receptors; cell-surface antigens & cell adhesion proteins; interleukins & interferons; cytoskeleton & motility proteins; and protein turnover. In a specific mouse array of interest, the spots are as listed in Table 2.

The mouse array of the subject invention finds use in a variety of different applications, where such applications include: profiling differential gene expression in transgenic knockout mice or other experimental mouse models; investigating processes such as embryo genesis and tumorigenesis; discovering potential therapeutic and diagnostic drug targets; and the like.

TABLE 2

D13473			
	MmRad51; yeast DNA repair protein Rad51 and E coli RecA homologue.	C6m	855-1199
D17630	Interleukin-8 receptor	E3h	664-1022
	Catenin alpha	E5m	1276-1594
D31788	BST-1; lymphocyte differentiation antigen CD38	B2h	674-1014
D31942	Oncostatin M	F3n	1017-1360
L05630	C5A receptor	E1g	841-1165
L07264	Heparin-binding EGF-like growth factor (Diphtheria toxin receptor)	F2d	258-673
U04807	Fms-related tyrosine kinase 3 Flt3/Flk2 ligand	C3i	46-418
	CD27; lymphocyte-specific NGF receptor family member	C2I	596-846
M28998	Fibroblast growth factor receptor Basic (b FGF-R)	E2c	200-583
M58288	Granulocyte colony - stimulatings factor receptor	E1)	251-529
	Growth/ difflerentiation factor 1 (GDF-1) (TGF- beta family)	F2b	2267-2566
	PKC-delta; protein kinase C delta type	B6g	1740-2011
	GA binding protein beta-2 chain	D3d	613-931
M83312	CD 40L receptor (TNF receptor family)	E1f	417-754
M83649	Fasl receptor (Fas antigen, Apo-1 antigen)	C3f	416-736
M86671	Interleukin 12 (p40) beta chain	F4n	652-963
M95200	Vascular endothelial growth factor (VEGF)	F4j	688-955
U03421	Interleukin 11 (adipogenesis inhibitory factor)	F4m	196-475
U14332	Interleukin 15	F5a	605-1057
U15159	LIMK; LIM serine/threonine kinase	B5I	1376-1699
U83628	DAD-1; defender against cell death 1	PEO	221-509
U25416	CD 30L receptor (Lymphocyte activation antigene CD 30. Ki-1 antigene)	C2m	135-435
U44725	Mast cell factor	F3i	79-417
	C-C chemokine receptor (Monocyte chemoattractant protein 1 receptor		
U56819	(MCP-1RA)	E1d	965-1262
X06381	Leukemia inhibitory factor (LIF) (cholinergic differentiation factor)	F3d	63-366
X52264	Intercellular adhesion molecule-1	E7i	1053-1385
89763	Interleukin-1 receptor type II	E2n	883-1134
X72305	Corticotropin releasing factor receptor	E1h	1411-1748
X72307	Hepatocyte growth factor (hepapoitein)	F2e	641-965
Z22703	Keratinocyte growth factor FGF-7	F3b	63-325
Z31663	Activin type I receptor	E1a	847-1130

TABLE 2 (CONT)

GenBank #	GenBank # Gene Name	Array Coordinate	Position
D01034	Transcription factor TF II D	B4j	291-556
9	20-1; Tight junction protein; discs-large family member, partially	704	
014340	FROCE excision renair protein DNA renair protein XP C	AZO	3/14-4001
D16306	chicos excisión repair protein, braz repair protein complementario Artas (cells (XPG)	Cef	1336-1639
L22472	Bax; Bcl-2 heterodimerization partner and homologue	C1g	172-534
	; CD28 antigen ligand 2, B7-2		
L25606	antigen; alternative CTLA4 counter-receptor	B2g	570-967
	NF2; Merlin (moesin-ezrin-radixin-like protein); shwannomin, murine		
L27105	neurofibromatosis type 2 susceptibility protein	A1i	2175-2400
M13945	Pim-1 proto-oncogene	A4a	2713-2930
M20157	Egr-1 Zn-linger regulatory protein	D2i	399-753
M25811	PKC-alpha; protein kinase C alpha type	B6e	1566-1924
M27129	CD44 antigen	E6e	789-1141
M31042	T-lymphocyte activated protein	D6h	285-606
M31131	Neuronal-cadherin (N-cadherin)	E7k	1212-1409
	ATP-dependent DNA helicase II 70 kDa subunit; thyroid Ku (p70/p80)		
M38700	autoantigen p70 subunit; p70 Ku)	C5h	274-632
M63660	G13; G-alpha-13 guanine nucleotide regulatory protein	B6n	2057-2377
M83380	Transcription factor ReIB	D7c	1456-1728
M84487	Vascular cell adhesion protein 1	E7m	984-1304
	ERCC3 DNA repair helicase; DNA-repair protein complementing XP-B cells		
S71186	(XPBC)	C6e	1147-1444
S76657	CRE-BP1; cAMP response element binding protein 1	Взі	412-748
U02887	XRCC1 DNA-repair protein, affecting ligation	C7n	900-1183
U53228	Nuclear hormone receptor ROR-ALPHA-1	DSi	368-675
U57311	14-3-3 protein eta	B7g	374-640
X56135	Prothymosin alpha	A7m	186-455
X57487	PAX-8 (paired box protein PAX 8)	DSI	680-1011
X58995	CamK IV; Ca2/calmodulin-dependent protein kinase IV (catalytic chain)	BSf	1269-1608
	ATP-dependent DNA helicase II 80 kDa subunit; thyroid Ku (p70/p80)		
X66323	autoantigen p80 subunit; p80 Ku)	C5i	565-875
X67812	Ret proto-oncogene (Papillary thyroid carcinoma-encoded protein)	A4f	2359-2680
0000	Nm23-M2; nucleoside diphosphate kinase B; metastasis-reducing protein;		
X68193	c-myc-related transcription factor	C4c	80-454

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
X97052	MAPKK6; MAP kinase kinase 6(dual specificity) (MKK6)	B6d	375-711
D17384	DNA polymerase alpha catalytic subunit (p180)	CSI	563-908
	Caspase-3; Nedd2 cysteine protease (positive regulator of programmed	;	
D28492	cell death ICH-1 homologue)	C1b	398-694
D50621	PSD-95/SAP90A	Ded	1512-1889
J04946	Angiotensin-converting enzyme (ACE) (clone ACE.5.)	F6f	850-1113
	Clusterin; complement lysis inhibitor; testosterone-repressed prostate		
L08235	message 2; apolipoprotein J; sulfated glycoprotein-2	C3b	515-744
L12721	Adipocyte differentiation-associated protein	D1c	404-709
121671	Epidermal growth factor receptor kinase substrate EPS8	D2k	1592-1873
L33768	Jak3 tyrosine-protein kinase; Janus kinase 3	BSj	3123-3426
L33779	Desmocollin 2	E6l	1317-1691
L47650	Stat6; signal transducer and activator of transcription 6; IL-4 Stat; STA6	B4g	2057-2411
M12056	Lymphocyte-specific tyrosine-protein kinase LCK	A5a	1205-1488
M22115	ERA-1 Protein (ERA-1-993)	D2I	723-1062
M26283	Homeo Box protein 2.1 (Hox-2.1)	D4a	647-884
M32309	Zinc finger X-chromosomal protein (ZFX)	D7n	2153-2554
M55512	WT1; Wilms tumor protein; tumor suppressor	A2c	1262-1563
M57422	Tristetraproline	B4k	262-504
M96823	Nucleobindin	D5j	80-357
M97013	PAX-5 (B cell specific transcription factor)	D6a	286-629
	IFN9R2; interferon-gamma receptor second (beta) chain; interferon gamma		
S69336	receptor accessory factor-1 (AF-1)	B3b	832-1089
S74227	Transcriptional enhancer factor 1 (TEF-1)	D7i	934-1233
U02079	Transcription factor NFAT 1, isoform alpha	D7a	1601-1910
U05252	DNA-binding protein SATB1	D2e	1101-1380
	CCHB3; calcium channel (voltage-gated; dihydropyridine-sensitive; L-type)		
U20372	beta-3 subunit)	B2c	351-639
	p57kip2; cdk-inhibitor kip2 (cyclin-dependent kinase inhibitor 18) member		-1
U20553	of the p21CIP1 Cdk inhibitor family, candidate tumor suppressor gene	A79	989-1272
U36203	snoN; ski-related oncogene	E2j	671-1006
X14759	Homeo Box protein 7.1 (Hox-7.1)	D4f	740-992
X14943	Neuronal cell surface protein F3	E71	1033-1311
X55123	GATA-3 transcription factor	D3f	858-1125
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TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
X57621	YB1 DNA binding protein	[70]	550-873
X58384	Dipeptidyl peptidase iv	E7i	61-294
X59421	Fii-1 ets-related proto-oncogene	A3b	267-623
X66224	RXR-beta cis-11-retinoic acid receptor	B4c	1225-1477
X78445	C3H cytochrome P450; Cyp1b1	B1j	295-593
X96859	Ubiquitin-conjugating enzyme, yeast Rad6 homologue; murine HR6B	C7k	51-392
Z27088	Relaxin	C4i	51-365
227410	Transcription factor LIM-1	D6m	1673-1934
D10061	DNA topoisomerase I (Top I)	C5m	1051-1357
D12513	DNA topoisomerase II (Top II)	CSn	520-870
D30687	GST Pi 1; glutathione S-transferase Pi 1; preadipocyte growth factor	C2d	62-369
J03958	Glutathione S-transferase A	C1n	54-311
J04696	Glutathione S-transferase Mu 1	C2b	13-263
L10656	c-Abl proto-oncogene	A4k	878-1145
M13071	A-Raf proto-oncogene	A3k	1042-1320
M17031	c-Src proto-oncogene	A4n	452-758
M35523	Retinoic acid binding protein II cellular (CRABP-II)	D6e	276-571
M83749	Cyclin D2 (G1/S-specific)	A6g	781-1074
U43844	Cyclin D3 (G1/S-specific)	A6h	484-790
S49542	5-Hydroxytryptamine receptor (Serotonin receptor type 2 (5HT2))	E4e	400-707
S78355	Cyclin D1 (G1/S-specific)	A6f	1858-2205
	Pur-alpha transcriptional activator; sequence-specific ssDNA-binding		
002098	protein	C7e	1082-1309
U27323	Cdc25a; cdc25M1; MPI1 (M-phase inducer phosphatase 1)	A7j	606-986
X07414	ERCC-1; DNA excision repair protein	C6d	189-484
X15842	c-rel proto-oncogene	A2m	1729-2064
X69618	Inhibin alpha subunit	F2g	810-1117
X76341	Glutathione reductase	C1m	115-377
X81581	Insulin-like growth factor binding protein-3 (IGFBP-3)	F2k	474-719
226580	Cyclin A (G2/M-specific)	A6a	701-1009
Z46845	Preproglucagon	A5i	172-531
	NF-kB p65; NF-kappa-B transcription factor p65 subunit; rel-related		
M61909	polypeptide	B4a	101-363
D11091	PKC-theta; protein kinase C theta type	B6h	658-957
D13867	VLA-3 alpha subunit	E7n	288-589

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
D17571	NADPH-cytochrome P450 reductase	C4a	326-605
D17584	Beta-protachykinin a	A5j	273-523
D30743	Wee1/p87; cdc2 tyrosine 15-kinase	A7h	1816-2159
D83966	Protein tyrosine phosphatase	C4g	1060-1429
J05205	Jun-D; c-jun-related transcription factor	A3g	737-964
L23423	Integrin alpha 7	E7e	2399-2713
L28177	Gadd45; growth arrest and DNA-damage-inducible protein	c3j	144-434
L35049	Bcl-xL apoptosis regulator (bcl-x long); Bcl-2 family member	C1j	641-906
X03919	N-myc proto-oncogene protein	A3j	3262-3450
M20473	cAMP-dependent protein kinase type I-beta regulatory chain	BSg	538-750
M21065	IRF1; interferon regulatory factor 1	B7k	1-233
M36830	HSP86; heat shock 86kD protein	B1d	255-551
	LFA1-alpha; integrin alpha L; leukocyte adhesion glycoprotein LFA-1 alpha		
M60778	chain; antigen CD11A (p180)	Взе	1838-2050
M88127	APC; Adenomatous Polyposis Coli protein	A1a	4127-4476
S93521	Cdc25b; cdc25M2; MPI2 (M-phase inducer phosphatase 2)	A7k	1893-2200
U03279	PI3-K p110; phosphatidylinositol 3-kinase catalytic subunit	B6j	1437-1723
003560	HSP27; heat shock 27kD protein 1	Bla	245-500
U05247	Csk; c-Src-kinase and negative regulator	B4n	645-984
	Fast; Fas antigen ligand; generalized lymphoproliferation disease gene		
U06948	(gld) in mice	C3g	168-488
U10871	MAPK; MAP kinase; p38	B5m	465-780
U19597	p19ink4; cdk4 and cdk6 inhibitor	A7d	228-516
U19617	Elf-1 Ets family transcription factor	D2j	1585-1902
U21050	CRAF1; TNF receptor (CD40 receptor) associated factor; TRAF-related	C3c	1225-1466
	SPI3; serpin; similar to human proteinase inhibitor 6 (placental thrombin	;	
025844	inhibitor) serine proteinase inhibitor	C4I	915-1230
	RIP cell death protein; Fas/APO-1 (CD95) interactor, contains death		
U25995	domain	C4j	1945-223
U29056	SLAP; src-like adapter protein; Eck receptor tyrosine kinase-associated	B5c	109-427
U43678	Atm; ataxia telangiectasia murine homologue	C5g	8989-9170
U51196	EB1 APC-binding protein	A1e	607-834
U51907	TANK; I-TRAF; TRAF family member associated NF-kB activator	B4h	135-437

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
	Caspase-11; ICH-3 cysteine protease; upstream regulator of ICE	C1a	352-686
U59883	MLH1 DNA mismatch repair protein; MutL homologue	Cek	1037-1278
X04480	Insulin-like growth factor-IA	F3a	183-406
X07640	Cell surface glycoprotein MAC-1 alpha subunit	E6j	1892-2179
X13664	N-ras proto-oncogene; transforming G-protein	A5e	548-857
X13945	L-myc proto-oncogene protein	A3h	5287-5590
X14951	CD18 antigen beta subunit (leukocyte adhesion LFA-1) (CD3, P150, 95)	E5n	1366-1706
X52191	c-Far proto-oncogene	A4m	1305-1538
X53176	Integrin alpha 4	E7b	2176-2449
X53532	PKC-beta; protein kinase C beta-II type	B6f	1712-2089
	HSP60; heat shock 60 kDa protein 1 (chaperonin, GroEL homologue);		
X53584	mitochondrial matrix protein P1	B1b	1432-1459
X57111	c-Cbl proto-oncogene (Adaptor protein)	A5b	858-1151
X59868	Cdc25 phosphatase; guanine nucleotide releasing protein	A7i	942-1276
	Ezrin; Villin 2; NF-2 (merlin) related filament/plasma membrane associated		
X60671	protein	A1f	1571-1812
X64713	Cyclin B1 (G2/M-specific)	A6c	1184-1447
X69902	Integrin alpha 6	E7d	261-611
X72395	5-Hydroxytryptamine (serotonin) receptor 3	E4j	1422-1711
X73573	Homeobox protein HOXD-3	D4h	141-362
X75888	Cyclin E (G1/S-specific)	A6i	799-1140
X76850	MAPKAPK-2; MAP kinase-activated protein kinase; MAPKAP kinase 2	B5n	719-987
X83971	Fra-2 (fos-related antigen 2)	A3d	617-844
X84311	Cyclin A1 (G2/M-specific)	A6b	656-916
	DCC; netrin receptor; immunoglobulin gene superfamily member; former		:
X85788	tumor suppressor protein candidate	A1d	4193-4508
	MHR23A; Rad23 UV excision repair protein homologue; xeroderma		
X92410	pigmentosum group C (XPC) repair complementing protein	Cei	613-955
	MHR23B; Rad23 UV excision repair protein homologue; xeroderma		
X92411	pigmentosum group C (XPC) repair complementing protein	Cej	542-807
Y00769	Integrin beta	E79	1990-2320
Z32767	MmRad52; yeast DNA repair protein Rad52 homologue	Cen	159-417
Z37110	Cyclin G (G2/M-specific)	A6k	300-619
D13458	Prostaglandin E2 receptor EP4 subtype	B3f	1146-1442
70000	Interleukin-5 receptor	E3f	1389-1739

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
J00380	Epidermal growth factor (EGF)	F1j	180-505
J04843	Erythropoietin receptor	E2a	1193-1377
J05149	Insulin receptor	E4a	653-1011
K01700	p53; tumor suppressor; DNA-binding protein	A11	1125-1517
L03529	Cf2r; coagulation factor II (thrombin) receptor	B2j	762-1154
L09562	PTPRG; protein-tyrosine phosphatase gamma	121	1248-1504
L10075	DNA-binding protein SMBP2	D2f	4790-5088
L12120	Interleukin-10 receptor	E3a	1762-2110
L20048	Interleukin-2 receptor gamma chain	E3c	1073-1313
L24755	Bone morphogenetic protein 1	F1b	2402-2676
L33406	Uromodulin	F4i	1809-2136
L34169	Thrombopoietin	F4e	652-954
M13177	Transforming growth factor beta	F4f	772-1075
M13926	Granulocyte colony- stimulating factor (G-CSF)	F2a	86-377
M14220	Neuroleukin	F3m	1110-1490
M14951	Insulin-like growth factor-2 (somatomedin A)	F2n	46-328
M15131	Interleukin 1 beta	F4k	827-1225
M16449	c-myb proto-oncogene protein	A2k	1212-1513
M16819	Tumor necrosis factor beta TNF-beta (Lymphotoxin-alpha)	F4h	461-805
M20658	Interleukin-1 receptor	C3u	2050-2410
X05010	CSF-1; M-CSF; colony stimulating factor-1	A5g	1268-1657
M27959	Interleukin-4 receptor (membrane-bound form)	E3e	2469-2705
M28233	Interferon-gamma receptor	E2m	1262-1550
M29697	Interleukin-7 receptor	E3g	701-1104
M34815	Gamma interferon induced monokine (MIG)	F1m	42-323
M37897	Interleukin 10	F4I	175-456
M57999	NF-kappa B binding subunit (nuclear factor) (TFDB5)	D5g	3122-3417
M59378	Tumor necrosis factor receptor 1; TNFR-1	CSd	1961-2376
M84607	PDGFRa; platelet-derived growth factor alpha-receptor	A4e	474-803
M84746	Interleukin-9 receptor	E3i	795-1086
M87039	iNOSI; nitric oxide synthase (inducible)	C3m	3178-3455
M89641	Interferon alpha-beta receptor	E2I	808-1120
M94087	Activating transcription factor 4 (mATF4)	D1b	416-769
S56660	Beta2-RAR; retinoic acid receptor beta-2	B3k	968-689
S67051	Tia-2 proto-oncogene	A4i	1843-2179

TABLE 2 (CONT)

ConBook #	Gene Name	Array Coordinate	Position
1100182	IGE-I-B alpha: insulin-like growth factor I receptor alpha subunit	C3I	489-885
	IGFR II; insulin-like growth factor receptor II, cation-independent mannose-		
U04710	6-P receptor, elevated in Wilms's tumor cells	C3k	707-1060
U06922	Stat3; APRF; acute phase response factor	B4e	1575-1910
U18542	Calcitonin receptor 1b	E3k	1375-1630
U32329	Endothelin b receptor [Ednrb]	E1i	279-695
U32330	Prepro-endothelin-3	F4c	703-1008
X04367	Pre-platelet-derived growth factor receptor	E2i	2336-2677
X04836	CD 4 receptor (T cell activation antigene)	E1e	1652-1877
X07962	Interleukin 7	F5d	241-496
X12531	Macrophage inflamatory protein	F3e	25-359
X14432	Thrombomodulin	F4d	1082-1365
X51975	Interleukin 6 (B cell differentiation factor)	F5c	1638-1898
X53779	Androgen receptor	E3j	2189-2491
X56848	Bone morphogenetic protein 4 (BMP-4) (TGF-beta family)	F1d	1275-1513
X57349	Transferrin receptor protein (p90, CD71)	Bah	654-1023
X57413	Transforming growth factor beta 2	F4g	2227-2541
X57497	Glutamate receptor, ionotropic AMPA 1	E5h	1290-1657
X57796	TNF 55; tumor necrosis factor 1 (55kd)	CSb	656-1022
X58876	Mdm2; p53-regulating protein	A1h	1364-1646
X61753	Transcription factor 1 for heat shock gene	Dei	203-570
X65453	CD40L; CD40 ligand	C2n	545-809
	c-Fms proto-oncogene (macrophage colony stimulating factor 1 (CSF-1)		
X68932	receptor)	A4b	2399-2686
X70472	B-myb proto-oncogene; myb-related protein B	A2f	2109-2456
X76654	Ear-2; v-erbA related proto-oncogene	AZn	1065-1376
X80764	Tie-1 tyrosine-protein kinase receptor	B3g	1425-1844
D10651	Glutamate receptor, ionotropic NMDA2B (epsilon 2)	E5j	506-786
D10217	Glutamate receptor, ionotropic NMDA2A (epsilon 1)	E5i	3966-4209
D10329	CD7 antigen	E6g	28-421
D00926	Transcription factor S -II (transcription elongation factor)	DZd	518-767
D12482	Basic Fibroblast growth factor (b- FGF)	F1a	290-620
D16250	Bone morphogenetic protein receptor	E1c	1454-1837
D17292	G-protein-coupled receptor	E2d	833-1115
D17407	Transcription factor SP2	D7g	734-1079

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
D29678	Cdk5; cyclin-dependent kinase 5	A6n	552-882
D25540	TGF-beta receptor type 1	E2k	1407-1629
D26077	Kinesin like protein KIF 3B	F6a	3519-3722
D29951	Kinesin family protein KIF1A	F5m	2553-2830
D38258	Fibroblast growth factor 9	F1K	91-379
D83698	Neuronal death protein	C4b	627-805
D84372	Syp; SH-PTP2; adaptor protein tyrosine phosphatase	BSe	1229-1543
J03168	Interferon regulatory factor 2 (IRF 2)	D4I	718-976
J02870	Lamimin receptor 1	E7j	368-675
D90176	NF-1B protein (transcription factor)	DSf	452-791
J03236	Jun-B; c-jun-related transcription factor	A3f	514-740
103520	Tissue plasminogen activator	F7e	622-1020
J03770	Homeo Box protein 4.2 (Hox-4.2)	D4e	565-945
J04113	Nur77 early response protein; thyrold hormone (TR3) receptor	C4d	825-1059
J04103	Ets-2 transcription factor	D3b	917-1281
J04115	c-Jun proto-oncogene (transcription factor AP-1 component)	A2i	951-1238
305609	Serine protease inhibitor homolog J6	F71	581-855
K01759	Nerve growth factor beta (beta-NGF)	F3I	642-901
L01640	Cdk4; cyclin-dependent kinase 4	A6m	230-616
K02582	Acetylcholine receptor delta submit	E4I	1400-1655
L02526	MAPKK1; MAP kinase kinase 3 (dual specificity) (MKK1)	Вба	1284-1583
L04662	GABA-A transporter 4	E5g	960-1341
L04663	GABA-A transporter 3	ESf	1010-1320
	Vegfr1; Vascular endothelial growth factor receptor 1 / Fms-related tyrosine		
L07297	kinase 1 (Flt1)	A4j	1144-1541
L10084	Adrenergic receptor, beta 1	E4m	404-772
L25890	Eph3 (Nuk) tyrosine-protein kinase receptor	B2k	2255-2491
L16953	MTJ1; DnaJ-like heat-shock protein from mouse tumor	B1e	1059-1384
L19622	TIMP-3 tissue inhibitor of metalloproteinases-3	F7n	274-592
L24563	Insulin receptor substrate-1 (IRS-1)	E4b	1027-1304
L13968	YY1 (UCRBP) transcriptional factor	D7k	1052-1292
L28095	Interleukin-converting enzyme (ICE)	F7a	30-269
L38847	Hepatoma transmembrane kinase ligand	F2f	927-1219
L36179	Voltage-gated sodium channel	B2f	4179-4505
L37296	Bad; heterodimeric partner for Bcl-XL and Bcl-2; promotes cell death	C1d	1079-1375

TABLE 2 (CONT)

GenRank #	Gene Name	Array Coordinate	Position
1.35236	Jnk stress-activated protein kinase (SAPK)	B5k	795-1032
M11686	Cytoskeletal epidermal keratin (18 human)	F5i	473-773
M11434	Nerve growth factor aloha (aloha-NGF)	F3k	294-494
M10937	Foidermal keratin (1 human)	F5k	326-683
M14537	Nicotinic acetylcholine receptor	ESK	1226-1568
M14757	MDR1: P-olycoprotein: multidrug resistance protein; efflux pump	B1g	1500-1886
M18934	CD2 antiden	E6a	354-602
M17192	Homeo Box protein 1.1 (Hox-1.1)	Dan	466-723
M19436	Fetal myosin alkali light chain	FSI	205-504
M25892	Interleukin 4	F5b	77-310
	Rb; pp105; Retinoblastoma susceptibility-associated protein (turnor		
M26391	suppressor gene; cell cycle regulator)	A1m	2036-2296
M28489	Rsk: ribosomal protein S6 kinase	B6i	1191-1436
M29464	Pletelet- derived growth factor (A chain) (PDGF- A)	F4b	152-425
M28698	Cytoskeletal epidermal keratin (19 human)	F5j	194-500
M29475	RAG-1: V(D)J recombination activating protein	C7g	2155-2404
M29855	Interleukin-3 receptor	E3d	1975-2254
M30642	K-fibroblast growth factor	F3c	309-577
M34381	Octamer binding transcription factor (Oct 3)	D5k	774-999
M33960	Plasminogen activator inhibitor	F7h	1096-1344
M33158	CD3 antigen, delta polypeptide	E6c	73-361
M34857	Homeo Box protein 2.5 (Hox-2.5)	D4c	11-277
M36829	HSP84: heat shock 84kD protein	B1c	342-366
M55617	Mast cell protease (MMCP) - 4	F7b	634-992
M61177	Frk1: extracellular signal-regulated kinase 1; p44; Ert2	B5h	115-373
	P13-K p85; phosphatidylinositol 3-kinase regulatory subunit;	,	
M60651	phosphoprotein p85; PDGF signaling pathway member	B6k	981-1260
	p58/GTA; galactosyltransferase associated protein kinase (cdc2-related		
M58633	protein kinase)	A7b	1022-1284
M64086	Serine protease inhibitor 2 (spi-2)	F7j	1499-1754
M64429	B-Raf proto-oncodene	A3i	1651-2036
M68513	Etk1 (Mek4; HEK) tyrosine-protein kinase receptor HEK	B2I	2681-2915
M64796	RAG-2: V(D)J recombination activating protein	C7h	671-944
M84324	Collagenase type IV	F6k	696-1040
M83336	Interleukin-6 receptor beta chain; membrane glycoprotein gp130	B3c	1423-1741

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
M76601	Alpha cardiac myosin heavy chain	F5e	2094-2391
M84819	Retinoic acid receptor RXR- gamma	J9Q	701-1082
M85078	Granulocyte-macrophage colony-stimulating factor receptor	E2e	904-1289
M86566	GABA-A receptor alpha-1 submit	ESd	1251-1606
M93428	Endothelial ligand for L-selectin (GLYCAM 1)	F1i	182-541
M95633	Integrin beta 7 subunit	E7h	2142-2423
U00478	DNAse I	Cec	665-871
U03184	Cortactin; protein tyrosine kinase substrate	B7h	426-653
U05672	Adenosine A2M2 receptor	C2g	491-735
U04674	DNA ligase I	CSj	1678-2054
U05671	Adenosine A1M receptor	C2f	302-673
U04443	Non-muscle myosin light chain 3	F6b	84-370
U06119	Cathepsin H	F6i	325-694
U06924	Stat1; signal transducer and activator of transcription	B4d	1749-2104
U09507	p21/Cip1/Waf1; cdk-inhibitor protein 1	A7e	9-403
	Cdk7; MO15; cyclin-dependent kinase 7 (homologue of Xenopus MO15		
U11822	cdk-activating kinase)	A7a	454-824
U10440	p27kip1; G1 cyclin-Cdk protein kinase inhibitor, p21-related	A7f	270-454
U10551	Gem; induced, immediate early protein; Ras family member	B7a	220-471
U12570	VHL; Von Hippel-Lindau tumor suppressor protein	A2b	885-1111
U12983	Cek 5 receptor protein tyrosine kinase ligand	F1g	1037-1287
U13705	Glutathione peroxidase (plasma protein); selenoprotein.	C11	766-1046
U14135	Integrin alpha 5 (CD51)	E7c	2170-2516
U14173	Ski proto-oncogene	A4g	707-1037
017698	Ablphilin-1 (abi-1) similar to HOXD3	D1a	351-585
U17162	BAG-1; bcl-2 binding protein with anti-cell death activity	C1e	17-334
	Shc transforming adaptor protein; Src homology 2 (SH2) protein, SHB-		
U15784	related	ASÍ	1220-1451
	MAPKK4; MAP kinase kinase 4; Jnk activating kinase 1; (JNKK1; SEK1;		
U18310	MKK4)	B6c	1380-1749
U19118	Franscription factor LRG - 21	Den	618-966
U19119	Interferon inducible protein 1	D4K	1342-1636
U19463	A20 zinc finger protein; apoptosis inhibitor	C2e	1952-2293
U19596	p18ink4; cdk4 and cdk6 inhibitor	A7c	16-284
U19799	I-kB (I-kappa B) beta	B3n	419-778

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
U24160	Dvl2; dishevelled-2 tissue polarity protein	B7i	1205-1578
U20532	Nuclear factor related to P45 NF-E2	DSh	1429-1759
U21011	MSH2 DNA mismatch repair protein; MutS homologue 2	C7a	2150-2490
U20238	GapIII; GTPase-activating protein	B7j	328-644
U25685	Syk tyrosine-protein kinase (activated p21cdc42Hs kinase (ack))	B5d	1235-1 524
	p107; RBL1; Retinoblastoma gene product-related protein p107 (cell cycle	-	
U27177	regulator)	A1j	1973-2365
U28724	PMS2 DNA mismatch repair protein; yeast PMS1 homolog 2	C7d	749-1013
U29173	Limphotoxin receptor (TNFR family)	E2g	1415-1668
U31625	BRCA1; Breast/ovarian cancer susceptibility locus 1 product	A1b	5126-5430
U33626	Pml; Murine homologue of the leukemia-associated PML gene	B4b	1667-2064
U34960	Transducin beta-2 subunit	B7e	515-834
U36277	I-kB (I-kappa B) alpha chain	B3m	541-823
U37522	TRAIL; TNF-related apoptosis inducing ligand; Apo-2 ligand	CSc	981-1288
	p130; Retinoblastoma gene product-related protein Rb2/p130 (cell cycle		
U36799	regulator)	A1k	970-1321
U36340	CACCC Box- binding protein BKLF	D1j	826-1065
U39643	FAF1; Fas-associated protein factor, apoptosis activator	C3e	423-681
U41671	Zinc finger transcription factor RU49	D7m	1229-1591
U42190	GTBP: G/T-mismatch binding protein; MSH6	C6g	1477-1769
U43144	PLC beta; phospholipase C beta 3	B6i	1933-2271
	Frizzled-3; Drosophila tissue polarity gene frizzled homologue 3;		-
U43205	dishevelled receptor	B2m	2037-2285
U43187	MAPKK3; MAP kinase kinase 3 (dual specificity) (MKK3, MEK3)	B6b	1436-1742
U43525	Myeloblastin; trypsin-chymotrypsin related serine protease	A7I	503-807
U47104	Zinc finger Kruppel type Zip 92	D7I	578-896
U44088	TDAG51; couples TCR signaling to Fas (CD95) expression	CSa	729-1042
U43788	POU domain, class 2, associated factor 1	Dec	610-884
U48853	Cas; Crk-associated substrate; focal adhesion kinase substrate	B4I	1982-2216
U49112	ALG-2; calcium binding protein required for programmed cell death	C2i	527-861
U49739	Unconventional myosin VI	F6e	3784-4021
U51037	Transcription factor CTCF (11 zinc fingers)	Dei	1625-1911
U53925	Transcription factor C 1	D6k	3895-4227

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
	Madr1: mSmad1: Mothers against dpp protein (Mad) murine homologue:		
U58992	TGF-beta signaling protein-1 (bsp-1); candidate tumor suppressor gene	A1g	238-476
U59746	Bci-W apoptosis regulator; Bci-2 family member	C1i	153-368
U60530	Mad related protein 2 (MADR2)	F3h	584-820
U62638	Cyclin C (G1-specific)	A6e	714-986
U63386	Mph-1 nuclear transcriptional repressor for hox genes	D5a	1621-1884
U66887	Rad50; DNA repair protein	C7f	1383-1707
U70324	Fyn proto-oncogene; Src family member	B5a	584-882
X01023	c-myc proto-oncogene protein	A2i	379-667
	c-Fos proto-oncogene; transcription factor AP-1 component. fos cellular		
V00727	oncogene	A2h	482-734
X06086	Cathepsin L	F6j	267-588
X04648	Glutamate receptor channel subunit gamma	E6n	41-408
X12616	c-Fes proto-oncogene	A4I	2342-2598
X12822	Cytotoxic cell protease 2 (B10)	F6I	439-686
X07439	Homeo Box protein 3.1 (Hox-3.1)	D4d	449-722
X13721	Homeo Box protein 2.4 (Hox-2.4)	D4b	1949-2284
X14897	Fos-B; c-fos-related protein fos B	A3c	920-1278
X16490	Plasminogen activator inhibitor-2	F7i	674-978
X51983	c-ErbA oncogene; thyroid hormone receptor.	A2g	400-675
X53337	Cathepsin D	F6h	587-894
X51438	Vimentin	F6d	868-1096
X53476	HMG-14 non histone chromosomal protein	D3m	643-1017
X53798	Macrophage inflamatory protein 2 alpha (MIP 2 alpha)	F3g	14-352
90695X	Bone morphogenetic protein 7 (BMP-7) (osteogenic protein 1)	F1e	670-971
X56959	Transcription factor SP1P (POUdomain transcription factor)	D7f	866-1128
X59252	Homeo Box protein 8 (Hox-8)	D4g	826-1132
X59927	Fibroblast growth factor receptor 4	E2b	2446-2820
X57277	Rac1 murine homologue	B7c	425-651
X60831	Transcription factor UBF	, , , , , ,	689-993
X61435	Kinesin heavy chain	F5n	1898-2182
X61800	CCAAT- Binding transcription factor (C/ EBP)	D1k	904-1150
X62622	TIMP-2 tissue inhibitor of metalloproteinases-2	F7m	1236-1468
X63190	Ets-related protein PEA 3	D3a	1702-2040

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
X64361	Vav; GDP-GTP exchange factor; proto-oncogene	87f	1083-1351
X63963	PAX-6 (paired box protein)	Deb	1081-1325
X66032	Cyclin B2 (G2/M·specific)	A6d	874-1236
	Chop10; murine homologue of Gadd153 (growth arrest and DNA-damage-		
X67083	inducible protein)	C3a	17-332
X67914	PD-1 possible cell death inducer; lg gene superfamily member	C4f	1481-1734
X69619	Inhibin beta A subunit (TGF beta family)	F2h	1064-1304
	Vegfr2; KDR/fik1 vascular endothelial growth factor tyrosine kinase		
X70842	receptor	B3j	1394-1721
X70296	Protease nexin 1 (PN-1)	F7d	746-985
X71327	MRE-binding transcription factor	DSb	552-916
X72711	Activator -1 140 KD subunit (replication factor C 140KD)	CSe	4137-4375
X72310	DP-1 (DRTF-polipeptide 1) cell cycle regulatory transcription factor	D2g	925-1305
X72230	5-Hydroxytryptamine (serotonin) receptor 1c	E4g	982-1314
X72795	Gelatinase B	F6n	599-954
X74351	XPAC; xeroderma pigmentosum group A correcting protein	C7m	447-669
X75427	Integrin alpha 2 (CD49b)	E7a	1595-1976
X77113	Growth/ diffferentiation factor 2 (GDF-2)	F2c	939-1329
X81582	Insulin-like growth factor binding protein-4 (IGFBP-4)	F2I	781-1140
X81579	Insulin-like growth factor binding protein-1 (IGFBP-1)	F2j	27-256
	IGFBP-2; insulin-like growth factor binding protein 2; autocrine and/or		
X81580	paracrine growth promoter	A5m	449-817
X81583	Insulin-like growth factor binding protein-5 (IGFBP-5)	F2m	461-824
X81584	Insulin-like growth factor binding protein -6 (IGFBP 6)	F2i	701-1039
X82327	A-myb proto-oncogene; myb-related protein A	A2e	1017-1334
X83536	Membrane type matrix matalloproteinase	F7c	877-1101
X87257	Elk-1 ets-related proto-oncogene	A3a	1498-1680
X86925	E2F-5 transcription factor	D2h	426-728
X90829	Lbx 1 transcription factor	D4n	1000-1306
X91144	P-selectin (glycoprotein ligand-1)	E5I	1095-1323
X91753	Transcription factor SEF2	D7e	755-1054
211974	Macrophage mannose receptor	E2h	807-1197
X95403		B7b	232-505
X98055	gation enzyme	C2c	14-298
X99063	Zyxin; LIM domain protein; alpha-actinin binding protein	B7n	1437-1812

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
Y00671	Met protooncogene	A4d	3646-3933
	c-Kit proto-oncogene (mast/stem cell growth factor receptor tyrosine		
Y00864	kinase)	A4c	2867-3181
Y07960	Transcription factor BARX1 (homeodian transcription factor)	D6j	723-973
X95346	PLC gamma; phospholipase C gamma	B6m	180-516
Z12604	Stromelysin-3; matrix metalloproteinase-11 (MMP-11)	C4n	1463-1806
Z14224	5-Hydroxytryptamine (serotonin) receptor 1e beta	E4h	530-774
215119	5-Hydroxytryptamine (serotonin) receptor 2c	E4i	588-940
219521	Low density lipoprotein receptor	E4d	1047-1324
Z23107	5-Hydroxytryptamine (serotonin) receptor 7	E4k	460-817
	c-Mpl; thrombopoietin receptor; hematopoietic growth factor receptor		
Z22649	superfamily member	A5k	1561-1772
Z21848	DNA-polymerase delta catalytic subunit	Ceb	1256-1600
Z29532	Follistatin	F11	764-1053
247766	Cyclin F (S/G2/M-specific)	A6j	2431-2708
Z36885	Ets-related protein Sap 1A	D3c	1267-1521
232815	Net; ets related transcription factor, activated by Ras	A3i	1211-1595
248538	Stat5a; mammary gland factor	B4f	2269-2628
	Hek2 murine homologue; Mdk5 mouse developmental kinase; Eph -related		
249086	tyrosine-protein kinase receptor	B2n	1702-1930
D26177	D-Factor/LIF receptor	E11	2376-2775
M13806	Cytoskeletal epidermal keratin (14 human)	F5h	108-469
M21019	R-ras protein, closely related to ras proto-oncogenes	B7d	215-555
M22959	Prolactin receptor PRLR2	E4c	1-328
M30903	Blk; B lymphocyte kinase; Src family member	C2j	1307-1672
M35590	Macrophage inflamatory protein 1 beta (Act 2)	F3f	119-445
M75716	Alpha-1 protease inhibitor 2	F7g	652-969
M92378	GABA-A transporter 1	E5e	1131-1416
M97017	Bone morphogenetic protein 8a (BMP-8a) (TGF-beta family)	F11	788-1139
M97200	Erythroid kruppel-like transcription factor	D2n	783-1171
M98339	GATA binding transcription factor (GATA-4)	D3e	81-379
M98547	Growth factor receptor	E2f	1701-2014
S72408	Crk adaptor protein	B4m	750-1027
U09419	Retinoid X receptor interacting protein (RIP 15)	D6g	1388-1682
U14752	Cek 7 receptor protein tyrosine kinase ligand	F1h	504-837

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
	C-C CKR-1; CCR-1; C-C chemokine receptor type 1, macrophage		
U2967B	inflammatory protein-1 alpha receptor; MIP-1alpha-R; RANTES-R	B2i	168-495
X13358	Glucocorticoid receptor form A	E3m	1527-1816
	Mothers against DPP protein (mad homolog Smad 1, transforming growth		
X83106	factor beta signaling protein)	F3j	464-728
Y00487	Hck tyrosine-protein kinase	B5b	1308-1563
AB000777	Photolyase/blue-light receptor homologue	C7c	1418-1737
D49482	Osp94 osmotic stress protein; APG-1; hsp70-related	B1f	1026-1266
D78645	Glucose regulated protein, 78kD; Grp78	B1m	167-411
	LCR-1; CXCR-4; CXC (SDF-1) chemokine receptor 4; HIV coreceptor		
D87747	(fusin); G protein-coupled receptor LCR1 homologue;	B3d	584-867
M23384	Glucose transporter-1, erythrocyte; Glut1	B2e	325-653
M80456	Int-3 proto-oncogene; NOTCH family member; NOTCH4	A5h	1846-2145
M94335	c-Akt proto-oncogene; Rac-alpha; proteine kinase B (PKB)	C2k	604-899
Y13231	Bak apoptosis regulator; Bcl-2 family member	C1f	1509-1786
U57324	PS-2; homologue of the Alzheimer's disease gene	C4h	437-783
U65594	BRCA2; Breast cancer susceptibility locus 2 product	A1c	649-922
U66058	DNA ligase III	C5k	2980-3205
U67321	Caspase-7; Lice2; ICE-LAP3 cysteine protease	C1c	1040-1280
075506	BID; apoptic death agonist	C1k	452-777
	WBP6; pSK-SRPK1; WW domain binding protein 6 serine kinase for SR		
U92456	splicing factors	B7m	482-774
U95826	Cyclin G2 (G2/M-specific)	A6I	408-688
X99018	Ung1; uracil-DNA glycosylase	C71	444-729
Y14019	Rab-3b ras-related protein	F6c	232-562
U28423	Inhibitor of the RNA-activated protein kinase, 58-kDa	B5i	180-487
U34259	Golgi 4-transmembrane spanning transporter; MTP	B2d	742-1060
U34920	ATP-binding casette 8; ABC8; hornolog of Drosophila white	B2b	1011-1319
U37720	CDC42 GTP-binding protein; G25K	F5g	1675-1982
U41751	Etoposide induced p53 responsive (EI24) mRNA	B11	1041-1296
U51866	Casein kinase II (alpha subunit)	A3n	1237-1517
U52945	TSG101 tumor susceptibility protein	A1n	446-713
U54705	Tumor suppressor maspin	A2a	251-507
097076	FLIP-L; apoptosis inhibitor; FLICE-like inhibitory protein	C3h	1476-1811
X63615	CamK II; Ca2+/calmodulin-dependent protein kinase II (beta subunit)	F5f	1951-2219

TABLE 2 (CONT)

# 1		Array Coordinate	Position
GenBank #	Cene Name	Station (Billy	10000
	Htk; Mdk2 mouse developmental kinase; Eph -related tyrosine-protein		4
Z49085	kinase receptor	B3a	2032-2365
D49921	Glial cell line-derived neurotrophic factor	Fin	236-539
106039	CD31 (Platelet endothelial cell adhesion molecule 1)	E6d	1172-1494
16928	CD22 antigen	E6i	2314-2645
139770	Gbx 2	D3g	1122-1395
M12302	Cytotoxic T tymphocyte-specific serine protease CCP I gene (CTLA-1)	F6m	585-830
M14222	Cathepsin B	F6g	382-729
M33324	Growth hormone receptor	E3n	1942-2240
M34563	CD28 (receptor for B71)	E6b	544-774
M38651	Estrogen receptor	E31	742-1013
S71251	Monotype chemoattractant protein 3	E1k	201-491
U03856	CD45 associated protein (CD 45-ap, LSM-1)	E6f	620-898
U11688	Orphan receptor	E1b	1686-1943
U17985	Cannabinoid receptor 1 (brain)	E4n	1091-1437
U43512	Dystroglycan 1	E6m	2267-2505
U46923	G-protein coupled receptor	ESc	350-671
X02389	Urokinase type plasminogen activator	F7f	1301-1538
X05719	CTLA-4 (immunoglobin superfamily member)	E6k	246-519
X56182	Myogenic factor 5	D5d	232-528
X62700	uPAR1; urokinase plasminogen activator surface receptor (CD87)	B3i	482-756
X69832	Serine protease inhibitor 2.4	F7k	621-927
X70298	SRY-box containing gene 4	D7b	34-311
L25602	Bone morphogenetic protein 2 (BMP-2) (TGF-beta family)	F1c	8372-8724
M10021 KO	M10021 [K022 [K02588] P-1-450; dioxin-inducible cytochrome P450	B2a	3729-4014
M16506	Bci-2; B cell lymphoma protein 2, apoptosis inhibitor	Cth	2125-2367
M34510	CD14 antigen	E6h	667-931
M81832	Somatostatin receptor 2	E3b	47-310
U19880	Dopamine receptor 4	Esb	907-1191
U21681	Cannabinoid receptor 2 (macrophage, CB2)	E5a	910-1262
U58533	Erf (Ets-related transcription factor)	D2m	1286-1613
211597	5-Hydroxytryptamine (serotonin) receptor 1b	E4f	1043-1355
D78382	Tob antiproliferative factor, interacts with p185erbB2	A7n	540-876
J03752	Glutathione S-transferase (microsomal)	C2a	185-428
L20331	Adenosine A3 receptor	C2h	182-382

TABLE 2 (CONT)

GenBank #	Gene Name	Array Coordinate	Position
U05341	p55cdc; cell division control protein 20	C4e	1061-1348
U12273	AP endonuclease; apurinic/apyrimidinic endonuclease (Apex)	CSf	1894-2150
X67735	Mas proto-oncogene (G-protein coupled receptor)	ASI	566-808
D26046	AT motif-binding factor ATBF1	D1d	9807-10112
D49474	HMG-box transcription factor from testis (MusSox17)	D3I	427-662
L03547.	Ikaros DNA binding protein	D4i	627-890
L12147	Early B cell factor (EBF)	D2a	750-1026
L12703	Engrailed protein (En-1) homolog	D2b	1323-1554
L12705	Engrailed protein (En-2) homolog	D2c	1626-1895
L21027	Transcription factor A10	B4i	499-806
L26507	Myocyte nuclear factor (MNF)	D5c	1203-1456
L36435	Basic domain/leucine zipper transcription factor	D1e	872-1073
M37163	Caudal type Homeobox 1 (Cdx1)	D1I	1040-1301
M58566	Butyrate response factor 1	D1i	768-1054
S53744	Brain specific transcription factor NURR-1	D1g	1548-1754
S68377	Brn-3.2 POU transcription factor	D1h	877-1237
S74520	Caudal type Homeobox 2 (Cdx2)	D1m	1085-1367
U01036	Erythroid transcription factor NF-E2	D2d	1-241
U20344	Gut-specific Kruppel-like factor GKLF	D3i	1558-1789
U25096	Kruppel-like factor LKLF	D4m	898-1193
U29086	Neuronal helix-loop-helix protein NEX-1	D5e	572-907
U36760	Brain factor 1 (Hfhbf1)	D1f	1080-1318
U41626	Split hand/foot gene	DSm	92-303
U42554	Sim transcription factor	D1n	2828-3066
U59876	Glial cells missing gene homolog (mGCM1)	D3h	727-1080
U62522	Sp4 zinc finger transcription factor	D4j	1704-1929
X61754	Heat shock transcription factor 2 (HSF 2)	D3j	1445-1640
X83974	RNA polymerase I termination factor TTF-1	A2j	3222-3433
L35949	Hepatocyte nuclear factor 3/forkhead homolog 8 (HFH-8)	D3k	913-1232
X94125	SRY-box containing gene 3 (Sox3)	DSn	212-443
D13759	Cot proto-oncogene	A3m	956-969
	HR21spA; protein involved in DNA double-strand break repair; PW29;		
D49429	calcium-binding protein	Ceh	103-434
1064407	MmLim15; RecA-like gene; DMC1 homologue; meiosis-specific	<u> </u>	504 704
101	Inclination recommends by organization of the property of the	200	10/-100

TABLE 2 (CONT)

		Array Coordinate	Position
Genbank #	FBn72 endoplasmic reticulum stress protein; protein disulfide isomerase-		
105186	related protein	B1k	1160-1470
\$50213	HMG1-related VDJ recombination signal binding protein	B1h	2263-2531
S65038	Gli oncogene: zinc finger transcription factor	A3e	104-505
1105245	Tiam-1 invasion inducing protein; GDP-GTP exchanger-related	A5n	4329-4628
116805	Sik: Src-related intestinal kinase	C4k	1246-1623
U28495	Lic proto-oncodene	A5d	853-1150
1140930	Oxidative stress-induced protein mRNA	B1n	1248-1561
U43900	STAM; signal transducing adaptor molecule	C4m	576-811
U46854	ShcC adaptor, Shc-related; brain-specific	C7i	246-601
U58987	MmMre11a putative endo/exonuclease	B1i	866-1204
X53068	PCNA; proliferating cell nuclear antigen; processivity factor	C7b	53-320
X81464	Translin: recombination hotspot binding protein	[C7j	205-431
X96618	PA6 stromal protein: RAG1 gene activator	C6a	442-749
1118342	Sky proto-oncogene (Tyro3; Rse; Dtk)	A4h	1927-2286
750013	H-ras proto-oncogene; transforming G-protein	A5c	1307-1544
1 47239	ERBB-2 receptor (c-neu, HER2 protein tyrosine kinase)	E1m	16-42
1 47240	ERBB-3 receptor	E1n	4-243
U22516	Placental ribonuclease inhibitor (Angiogenin)	F4a	512-766
L00923	myosin I	G13	2578-2921
U459777	Ca2+ binding protein, Cab45	G20	597-1082
M10624	murine ornithine decarboxylase	G14	865-1252
X51703	ubiauitin	G5	123-547
100423	Hypoxantine-quanine phosphoribosyltransferase	G7	301-751
D78647	phospholipase A2	G6	446-813
131609	ribosomal protein S29	G21	5-244
M325999	glyceraldehyde-3-phosphate dehydrogenase	G12	765-1016
M12481	heta-actin	G19	25-564

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Cancer Array

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In the cancer arrays of the subject invention, the polynucleotide probe compositions on the array correspond to those genes which are associated, e.g. play a role in, cellular proliferative diseases, particularly cancer, where human genes are of particular interest in many embodiments. Types of genes that are typically represented on a cancer array of the subject invention include: oncogenes, tumor suppressors, cell cycle regulators, genome plasticity genes, apoptosis genes, cell differentiation genes, regulators of tumor host interaction and metastasis, such as extracellular matrix proteins, cell adhesion receptors, molecules that control cell invasion and motility, and genes associated with angiogenesis.

In certain embodiments, of particular interest is an array having the following types of genes represented on its surface: cell cycle/growth regulators; apoptosis; growth factors/cytokines; oncogenes/tumor suppressors; cell adhesion, motility and invasion; invasion regulators; GTP ases and their regulators; cadherins; intermediate filament markers; receptors; cell fate/development regulators; DNA damage/response/repair/ recombination; and angiogenesis regulators. In a specific cancer array of interest, the spots are as listed in Table 3.

The cancer array finds use in a variety of applications, including: monitoring cellular responses to therapeutic compounds; comparing expression profiles of tumors at different developmental stages; developing diagnostic tools for distinguishing closely related tumors; and the like.

In the following Table 3, as well as preceding Tables 1 and 2, the "position" coordinate refers to the actual nucleotide residues of the listed gene that are represented on the array.

TABLE 3

	# Juo	Array Coordinate Position	Position
Cell Cycle/Growth Hegulators	Gelibalin #	Signal Cook (Billy	
QUADRANT A			
CELL DIVISION CONTROL PROTEIN 2 HOMOLOG (EC 2.7.1) (P34	X05360	A1a	655-886
CELL DIVISION PROTEIN KINASE 2 (EC 2.7.1) (P33 PROTEIN	M68520	A1b	1774-2180
KINASE)	X66357	A1c	216-882
CELL DIVISION PROTEIN KINASE 4 (EC 2.7.1.) (PSK-J3)	M14505	A1d	372-693
CELL DIVISION PROTEIN KINASE 5 (EC 2.7.1.) (TAU PROTEIN KINASE II CATALYTIC SUBUNIT) (TPKII CATALYTIC SUBUNIT)	X66364	\ \ \	469.767
(KINASE PSSALRE).	X66365	Alf	315-663
CELL DIVISION PROTEIN NITAGE 7 (EC 2.7.1-) (CDK-ACTIVATING	1.20320		
KINASE) (CAN) (39 ND PROTEIN KINASE) (1 33 INC. 3) (CITY) (CITY)		A1g	89-305
CYCLIN-DEPENDENT KINASE 5 ACTIVATOR ISOFORM P391 PRECIRSOR (CDK5 ACTIVATOR) (P391).	U34051	Ath	763-1-62
CYCLIN-DEPENDENT KINASE 5 ACTIVATOR PRECURSOR (CDK5	X80343		
ACTIVATOR) (TAU PROTEIN KINASE II 23 KD SUBUNII) (TPKII DEGIII ATORY SUBLINID (P23) (P25) (P35).	•	A1i	551-941
ASEA: WEHASE INDICER PHOSPHATASE 1 (EC 3.1.3.48)	M81933	A1j	1632-1978
cdc25B; M-PHASE INDUCER PHOSPHATASE 2 (EC 3.1.3.48).	M81934; [S78187]	441.	2036 3602
(CDC25Hu2)	100707	AIK	331.623
cdc25C; M-PHASE INDUCER PHOSPHATASE 3 (EC 3.1.3.48).	M34005	A1m	144-459
CLK-1	1 29216	A1n	1106-1356
CLN-2	129220	A2a	551-1002
SEBINE THREONING-PROTEIN KINASE KKIALRE	X66358	A2b	276-461
SERINE/THREONINE-PROTEIN KINASE PCTAIRE-1	X66363	A2c	1114-1434
SERINE/THREONINE-PROTEIN KINASE PCTAIRE-2	X66360	A2d	954-1250
SERINE/THREONINE PROTEIN KINASE PCTAIRE-3	X66362	A2e	549-911
SEBINE/THREONINE PROTEIN KINASE PITALRE	L25676	A2f	367-635
CDC2-RELATED PROTEIN KINASE CHED	M80629	A2g	1388-1548
CDC2-RFI ATED KINASE PISSLRE	L33264	A2h	454-755
CYCLINA	X51688	A2i	876-1218
CYCLIN B1 G2/MITOTIC-SPECIFIC	M25753	A2j	979-1311
CYCLING 61/S:SPECIFIC	M74091	A2k	6670-7326
CYCLIN D1 (CYCLIN PRAD1) (BCL-1 ONCOGENE)	X59798; [M64349]	A2I	3427-3784
CYCLIN D2	D13639 [M90813]	A2m	3932-4284
CYCLIN D3	M92287	A2n	537-894

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
	M73812	A3a	1295-1658
CYCLIN G1	U47413 [L49504]	A3b	755-1035
CYCLIN G2	U47414 [L49506]	A3c	989-1254
CYCLIN H	U11791 [U12685]	A3d	717-1026
	U09579; [L25610]		
DIFFERENTIATION ASSOCIATED PROTEIN 6) (MDA-6) (P21) (CDK-			
INTERACTING PROTEIN 1) (CIP1) (WAF1) (CDKN1A) (CDKN1) (SDI1)			
		A3e	1745-2063
CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT INKINASE INHIBITOR P57) (P57KIP2)	U22398	A3f	1048-1316
CYCLIN-DEPENDENT KINASE 4 INHIBITOR A (CDK4I) (P16-INK4) (P16- L27211	127211		0101-0101
INK4A) (MULTIPLE TUMOR SUPPRESSOR 1) (MTS1). (CDKN2A)		A3g	482-836
CYCLIN-DEPENDENT KINASE 4 INHIBITOR B (P14-INK4B) (P15-INK4B) U17075, [L36844]	U17075; [L36844]		
		A3h	116-462
<u>D</u> .	U40343; [U20498]	A3i	750-952
	U10564	A3j	1259-1502
THREONINE-PROTEIN KINASE PLK (EC 2.7.1) (PLK-1)	U01038		
		A3k	1330-3233
	U38545	A3I	2862-3961
	D63878	A3m	381-675
MOLOG	S72008	A3n	66-379
	U00001	A4a	870-3474
-CONJUGATING ENZYME E2-CDC34	L22005	A4b	249-550
	U18291	A4c	45-378
	U63131	A4d	519-1464
	U77949	A4e	216-447
	X60188		
(INSULIN-STIMULATED MAPZ KINASE) (MAP KINASE 1) (MAPK 1) (P44 JERK1) (ERT2) (P44-MAPK) (MICROTUBULE-ASSOCIATED PROTFIN-2			
KINASE).		A4i	754-1094
(INASE 3 (EC 2.7.1) (ERK3)	X80692		
(MAP KINASE ISOFORM P97) (P97-MAPK).		A4g	806-1267
EXTRACELLULAR SIGNAL-REGULATED KINASE 4 (EC 2.7.1) (ERK4) X59727 MAP KINASE ISOFORM P63\ (P63-MAPK)	X59727	441	7000 0100
(INASE 5 (EC 2.7.1) (FBK5)	19597B		20/0-2334
(ERK4) (BMK1 KINASE)		A4i	1010-1267
EXTRACELLULAR SIGNAL-REGULATED KINASE 6 (EC 2.7.1) (ERK6) X79483	X79483		
(EHKS)		A4j	530-831

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
MITOGÉN-ACTIVATED PROTEIN KINASE P38 (EC 2.7.1) (MAP	L35253; [L35263]		
KINASE P38) (CYTOKINE SUPPRESSIVE ANTI-INFLAMMATORY DRUG			
BINDING PROTEIN) (CSAID BINDING PROTEIN) (CSBP) (MAX-		A4k	925-1204
INTERACTING PROTEIN 2) (MAP NINASE MAIZ).	196318		
STRESS-ACTIVATED FIGURING MINAGE SING (EC. 2.7.1.7) (C. 301.1) TERMINAT KINASE 1) (JNK-46)			952-1263
STRESS-ACTIVATED PROTEIN KINASE JNK2 (EC 2.7.1) (C-JUN N-	L31951		
TERMINAL KINASE 2) (JNK-55).		A4m	638-1000
STRESS-ACTIVATED PROTEIN KINASE JNK3 (EC 2.7.1) (C-JUN N-	U34819; [U07620]		
TERMINAL KINASE 3) (JNK3) (MAP KINASE P49 3F12).		A4n	1018-1413
DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE	U25265		
5 (EC 2.7.1) (MAP KINASE KINASE 5) (MAPKK 5) (MAPK/ERK KINASE		750	620.847
5).		ADA	053-047
DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE	L05624		·
1 (EC.2.7.1) (MAR NIKASE NIKASE 1) (MARTIN 1) (EIII) NOTION (INTERNATIONAL INTERNATIONAL INTERNATI		A5b	842-1217
DIA COCCEDENT MITOGEN ACTIVATED PROTEIN KINASE KINASE	139657		
DOAL STECHTOIL MILOGENACION ED THOUSEN WEST			
6 (SAPKK3)		A5c	1060-1389
MFK KINASE 3	U78876	A5d	1195-1453
PCNA (CYCLIN)	M15796; [J04718]	ASe	157-436
INIA	U49070	ASf	624-1075
RBP1(RETINOBLASTOMA-BINDING PROTEIN)	S57153; S57160	A5g	2676-2889
E2F-1 pRB-binding protein	W96577	A5h	899-1595
E2F-3	Y10479	A5i .	698-897
E2F-5	U15642	A5j	645-922
E2F-related transcription factor (DP-1)	L23959	A5k	935-1186
DP2 (Humdp2), dimerization partner of E2F	U18422	A5I	1603-1838
RBQ-3	X85134	A5m	359-603
GROWTH-ARREST-SPECIFIC PROTEIN 1 (GAS-1).	L13698	A5n	1550-1701
growth inhibitor p33ING1 (ING1)	AF001954	A6a	722-983
Abl interactor 2 (Abi-2) + Abl binding protein 3 (AbIBP3) [ArgBPIB]	1	A6b	1049-1203
GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2 (GRB2 ADAPTOR	L29511; [M96995]		
PROTEIN) (ASH PROTEIN).		A6c	355-573
GRB-IR / GRB10	U69276	A6d	358-1155
RAF ONCOGENE	X03484	A6e	1704-1989
raf,b-	M95712	A6f	866-1144
Jun B TRANSACTIVATOR	M29039	A6g	1197-1442
N-mvc	M13228	A6h	761-1188

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate	Position
C-myc binding protein	D89667	A6i	218-490
INTERMEDIATE FILAMENT MARKERS			
	Z29074; [S69510]	A6j	652-1781
KERATIN, TYPE I CYTOSKELETAL 10 (CYTOKERATIN 10) (K10) (CK 10)	M19156	A6k	295-497
Γ	D78367	AGI	455-624
	X52426; X07696; X62571		
13) +KERATIN, TYPE I CYTOSKELETAL 15 (CYTOKERATIN 15) (K15)			
(CK 15) +KEHAIIN, IYPE I CY I OSKELE I AL 17 (CY I OKEHAIIN 17)		Δ6m	383-1001
KERATIN, TYPE I CYTOSKELETAL 14 (CYTOKERATIN 14)(K14) (CK 14) J00124	J00124		
		A6n	339-839
KERATIN, TYPE I CYTOSKELETAL 16 (CYTOKERATIN 16)(K16) (CK	M21772; M20336	A7a	32-522
LETAL 18 (CYTOKERATIN 18) (K18) (CK	M26326		
18)		A7b	706-971
KERATIN, TYPE I CYTOSKELETAL 19 (CYTOKERATIN 19) (K19) (CK 19).	Y00503	A7c	726-1124
KERATIN, TYPE II CYTOSKELETAL 1 (CYTOKERATIN 1) (K1) (CK 1) (67 M98776 KD CYTOKERATIN) (HAIR ALPHA PROTEIN)	M98776	A7d	894-1459
KERATIN, TYPE II CYTOSKELETAL 2 ORAL (CYTOKERATIN 2P) (K2P) M99063 ICK 2P)	E9066W	A7e	2167-2455
KERATÍN, TYPE II CYTOSKELETAL 2 EPIDERMAL (CYTOKERATÍN 2E) M99061 [S43646] (K2E) (CK 2E)	M99061 [S43646]	A7!	1091-1450
KERATIN, TYPE II CYTOSKELETAL 4 (CYTOKERATIN 4) (K4) (CK4)		A7g	66-404
KERATIN, TYPE II CYTOSKELETAL 5 (CYTOKERATIN 5) (KS) (CK 5) (58 M21389 KD CYTOKERATIN)	M21389	A7h	93-682
KERATIN, TYPE II CYTOSKELETAL 6 (CYTOKERATIN 6A) (CK 6A)	J00269; V01516; L42592;		
(K6A KEHATIN) +(CY IOKEHATIN 6B) (CK 6B) (K6B KEHATIN) + (CYTOKERATIN 6C) (CK 6C) (K6C KERATIN) + (CYTOKERATIN 6D)	L00205; L42601; L42610; L42611; L42612		
(CK 6D) (K6D KERATIN) + (CYTOKERATIN 6E) (CK 6E) (K6E KERATIN) + (CYTOKERATIN 6F)		A7i	089-889
KERATIN, TYPE II CYTOSKELETAL 6B (CYTOKERATIN 6B) (CK 6B)	L42592; L00205		
			275-414
	X03212		1154-1430
KERATIN, TYPE II CYTOSKELETAL 8 (CYTOKERATIN 8) (K8) (CK 8)	M34225	A7I	1190-1474
VIMENTIN	X56134 [M14144]	A7m	460-740
DESMIN	U59167	A7n	1063-1364

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
QUADRANT B			
APOPTOSIS			
	M14745		5078-5382
RdP and n53 binding protein Bbo/53BP2 (BBP/53BP2)	U58334		3129-3376
BAX	. 22474		227-478
APOPTOSIS REGULATOR BCL-W	U59747	B1d	121-403
ELL DIFFERENTIATION PROTEIN	L08246	B1e	276-269
RCL 2-RELATED PROTEIN A1 (BFL-1 PROTEIN) (HEMOPOIETIC-	U29680		
SPECIFIC EARLY RESPONSE PROTEIN) (GRS PROTEIN)		B1f	64-293
BCL-2 INTERACTING KILLER (APOPTOSIS INDUCER NBK) (BP4)	X89986; [U34584]	B1g	935-1200
BECL'S (APOPTOGOUS ANTAGONIST/KILLER (APOPTOSIS	U23765; [U16812; U16811: X84213]	B1h	1371-1661
DAN DECTEIN (RCI -2 RINDING COMPONENT 6)	U66879	81i	408-749
BCL-2 BINDING ATHANOGENE-1 (BAG-1) (GLUCOCORTICOID	S83171; [Z35491]	81i	511-830
regine Abronine protein kingse NIK' binds specifically to TRAF2	Y10256		3776-4036
Casher, a FADD, and caspase-related inducer of apoptosis [CASH-alpha+ AF010127[Y14039;	AF010127[Y14039;		
CASH-betal (FLAME-1) (FLICE-like inhibitory protein)	Y14040]	B11	363-787
death domain containing protein CRADD, apoptotic adaptor molecule for	U84388	8	369-604
caspase-2 and Fast/INF receptor-interacting protein nir	141690	B1n	1009-1313
Interestion is associated process (T. 1922)	U25994; [U50062]	B2a	848-1123
DAXX a FAS-binding protein that activates JNK and apoptosis	AF015956	B2b	804-1030
Ano-2 ligand (TNF-related apoptosis inducing ligand TRAIL)	U57059	B2c	211-616
TRAF-INTERACTING PROTEIN I-TRAF (TRAF family member-associated U59863; [U63830]	<u> </u>	B2d	674-887
TRAFF	U69108	B2e	1318-1694
TRAF6	U78798; [L81153]	B2f	1689-1961
TRAE interacting protein (TRIP)	U77845	B2g	154-387
tumor necrosis factor type 2 receptor associated protein (TRAP3)	U12597		1207-1566
CD40 RECEPTOR ASSOCIATED FACTOR 1 (CRAF1) (CAP-1), (LMP1	U21092; [U15637; L38509;		
associated protein)	U19260]	B2i	980-1322
INHIBITOR OF APOPTOSIS PROTEIN 1 (HIAP1) (HIAP-1) (C-IAP2)	U45878; [U37546]		···
(AP1) (MHC).		B2j	1444-1848

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
INHIBITOR OF APOPTOSIS PROTEIN 2 (HIAP2) (HIAP-2) (G-IAP1) (TNFR2- TRAF SIGNALLING COMPLEX PROTEIN 2) (IAP HOMOLOG B)	U45879; [U37547]		
(IAP2) (MIHB).		B2k	266-621
X-LINKED INHIBITOR OF APOPTOSIS PROTEIN (X-LINKED IAP) (IAP-	U45880; [U32974]	. ICB	2000, 2263
ossachangedent cell growth requisitor CGB19	166469	R2m	28.301
pot deported to a grant recentor	190875	R2n	290-548
IICE INTEREST TOWNERTING ENZYME (PAS) (CASDASE I 113600: MAZEOT YESOTO)	113690- [M87507: Y65019]		200
(ICE) (INTERPORTED TO CONTRACT	o 15659, [mor 507, A05013]	B3a	5078-5282
(CASPASE-2) (ICH-1L) (ICH-1S)	U13021; [U13022]	B3b	851-1218
APOPAIN PRECURSOR (EC 3.4.22.) (CYSTEINE PROTEASE CPP32)	U13737		
(TAMA PHOLEIN) (CASPASE-S) (CPPSZ) (TAMA PHOLEIN) (CASPASE-)		830	2007-2434
ICH-2 PROTEASE PRECURSOR (EC 3.4.22) (TX PROTEASE) (ICEREL 1 U28014; U28015	U28014; U28015		
II) (CASPASE-4) + CASPASE-5 PRECURSOR (EC 3.4.22) (ICH-3			
PROTEASE) (TY PROTEASE (ICEREL-III).		B3d	763-11-07
CASPASE-6 PRECURSOR (EC 3.4.22) (APOPTOTIC PROTEASE MCH-U20537; U20536	U20537; U20536		
2) isoform beta + isoform alpha		ВЗе	387-697
CASPASE-7 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC	U37448		
PROTEASE 3) (ICE-LAP3) (APOPTOTIC PROTEASE MCH-3) (CMH-1)			
(Lice2)		B3f	1042-1413
CASPASE-8 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC PROTEASE 5) (MORTI-ASSOCIATED CED-3 HOMOLOG) (MACH)	U60520; U58143; X98172; X98173; X98174; AF00962		
(FADD-HOMOLOGOUS ICE/CED-3-LIKE PROTEASE) (FADD-LIKE ICE)			
(FLICE) (APOPTOTIC CYSTEINE PROTEASE) (APOPTOTIC			
PROTEASE MCH-5) (CAP4) (CASP8) (MCH5) isof		B3g	1327-1607
CASPASE-8 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC DEDITEASE 5) (MOBITA ASSOCIATED CED 3 HOMOLOG) (MACH)	U60520; U58143; X98172;		
(FADD-HOMOLOGOUS ICE/CED-3-LIKE PROTEASE) (FADD-LIKE ICE)	AF00962;X98176; X98175;		
(FLICE) (APOPTOTIC CYSTEINE PROTEASE) (APOPTOTIC	X98177; X98178		
PROTEASE MCH-5) (CAP4) (CASP8) (MCH5) isof	-	B3h	475-954
CASPASE-9 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC	U56390; [U60521]		
PROTEASE 6) (ICE-LAP6) (APOPTOTIC PROTEASE MCH-6)		B3i	986-1289
ICE-LIKE APOPTOTIC PROTEASE 4 PRECURSOR (EC 3.4.22)	U60519		
(APUPIULIC PHUIEASE MCH-4) (CASPASE-10)		B3j	2276-2690
DEATH-ASSOCIATED PROTEIN 3 (DAP-3) (ionizing radiation resistance conferring protein)	U18321; [X83544]	B3k	856-1114
DEATH-ASSOCIATED PROTEIN KINASE 1 (EC 2.7.1) (DAP KINASE 1). X76104	X76104	100	7000
		DOI:	1388-2321

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate	Position
FAST) phosphorvlates TIA-1	X86779	B3m	865-1239
	S78085	B3n	406-694
0.1	270519	B4a	1493-1887
GEN LIGAND (APOPTOSIS ANTIGEN LIGAND) (APTL)	D38122; [U08137]	B4b	1400-1782
WSL-LR, WSL-S1, WSL-S2 + TRAMP (Apo-3) (DDR3)	Y09392; [U75380;U74611;		
	U83597]	B4c	1407-1671
Akt1 (rac protein kinase alpha, protein kinase B, c-Akt)	M63167	B4d	
	M77198; [M95936]	B4e	1867-2099
	U69611	B4f	1540-1746
	AF016268	B4g	273-552
apoptosis gene/Bcl-2 homolog	S82185	B4h	351-995
seven in absentia homolog	U63295	B4i	239-523
BATS1	U37688	B4j	1247-1367
DNA fragmentation factor-45	U91985	B4k	485-1592
secreted apoptosis related protein 1	AF017986	B4I	189-974
secreted apontosis related protein 3 (SARP3)	AF017988	B4m	702-841
amontosis-related protein TFAR15 (TFAR15)	AF022385	B4n	365-520
celmodulin dependent phosphodiesterase PDE1B1	U56976	B5a	414-549
olutathione-S-transferase homolog	U90313	BSb	97-837
CD27BP (Siva)	U82938	B5c	406-625
chromosome segregation gene homolog CAS	U33286	B5d	674-1247
apoptosis inhibitor survivin	U75285	B5e	386-720
n53 induced protein	AF010310 AF010311	B5f	29-771
ping (PIG3)	AF010309	B5g	398-1223
Pig7 (PIG7)	AF010312	BSh	173-322
Pig10 (PIG10)	AF010314	B5i	437-1623
Pig11 (PIG11)	AF010315	B5j	748-1304
Pig12 (PIG12)	AF010316	BSK	97-531
GTP-binding protein (rhoA)	L25080	BSI	290-572
cdc42 homolog (G25K) [brain isoform + placental isoform]	M35543; [M57298]	B5m	321-468
ONCOGENES/TUMOR SUPPRESSORS			
C-FMS PROTO ONCOGENE	X03663	B5n	2568-2880
C-fos	K00650	B6a	2949-3181
± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ± ±	X06182	B6b	1981-2375
PROTO-ONCOGENE TYROSINE-PROTEIN KINASE SRC (EC 2.7.1.112)	HT2291; [K03214; X03996]	B6c	893-1189
PROTO-ONCOGENE TYROSINE-PROTEIN KINASE FGR (EC 2.7.1.112) M19722	M19722	P9G	521-856

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #		Position
ASH2	U04045; [L47583]	B6e	1496-2178
(mutS - ALPHA 160 KD TEIN) (GTBP) (GTMBP)	U54777		
		B6f	591-1100
ONCOGENE	M54968	B6g	352-604
	J02958	B6h	932-1242
	M14694; [M14695]	B6i	690-964
AST CANCER TYPE 2 SUSCEPTIBILITY PROTEIN	U43746	B6j	10056-10346
BRCA1-ASSOCIATED RING DOMAIN PROTEIN	U76638	B6k	1493-1801
+ MDM2-A (GB:	Z12020; [M92424]	198	920-1232
USSIBB) + MUME:C (GD: USSES)	AF007111	-	405-681
lated protein	Y11416	B6n	627-993
	X74594	B7a	951-1213
	X74262	B7b	605-974
BBP2 retinohlastoma hinding protein	S66431	B7c	2339-2642
	X85133	B7d	1701-1930
PROTO-ONCOGENE TYROSINE-PROTEIN KINASE RECEPTOR RET	M31213; [M57464]		
PRECURSOR (EC 2.7.1.112) (C-RE1), trapillaly lilyloid calcilloina-		B7e	2285-2631
Beticoblastoma suscentibility (RB1 retinoblastoma-assoc)	M15400	B7f	2839-3101
SKY (DTK) (TYRO3) (RSE)	D17517	B7g	2132-2597
VES VES	M15990	B7h	1325-1676
TYROSINE-PROTEIN KINASE BTK (EC 2.7.1.112) (BRUTON'S	U10087 X58957		
TYROSINE KINASEJ(AGAMMAGLOBOLINAEMIAT TROGINE KINASE) VATK) (R.CELL PROGENITOR KINASE) (BPK) (BTK) (AGMX1)		B7i	380-1430
TYPOSINE-PROTEIN KINASE ABL2 (EC 2.7.1.112) (TYPOSINE KINASE M35296 ABG) (ABL1)	M35296	B7j	493-1656
TYROSINE-PROTEIN KINASE ZAP-70 (EC 2.7.1.112) (70 KD ZETA- ASSOCIATED PROTEIN) (ZAP70)	L05148	B7k	1-584
SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 1- ALPHA/BETA (TRANSCRIPTION FACTOR ISGF-3 COMPONENTS	M97935	Ē	620 1276
P91/P84) (STAT1) SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 2	U18671 M97934	1/0	0.501-000
(P113) (STAT2)		B7m	1105-1480
SIGNAL TRANSDUCER AND TRANSCRIPTION ACTIVATOR 5B (STATSB)	U47686	B7n	831-1135
OUADBANT C		-	

TABLE 3 (CONT)

CONTRINGENORMENT U35835; [U47077] C1a C1a C1b C1	Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
PERUDEAT PROTEIN KINASE (DNA-PK) + DNA DEPENDENT U3893; [U47077] C1a	DNA DAMAGE RESPONSE/REPAIR/RECOMBINATION			
TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TELANGICCTASIA (ATM) TAGEN PROTEIN POJ (70 KD SUBUNIT (LUPUS KU NTIGEN PROTEIN POJ (70 KD SUBUNIT (LUPUS KU NTOSUM GROUP EACTON IV) TELANGIA (70 KD SUBUNIT (LUPUS KU NTOSUM GROUP EMENTING PROTEIN) TOON REPAIR PROTEIN RADS HOMOLOG TOTO ON REPAIR PROTEIN RADS HOMOLOG NTOSUM GROUP EMENTING PROTEIN) (DNA ON REPAIR PROTEIN ERCC.) (BASA THANSCRIPTION R 2 68 KD SUBUNIT) (STE2-969) (TFIH 69 KD SUBUNIT) EPAIR PROTEIN COMPLEMENTING PROTEIN) (DNA ON REPAIR PROTEIN ERCC.) EPAIR PROTEIN COMPLEMENTING PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN COMPLEMENTING PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN COMPLEMENTING PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN COMPLEMENTING PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN (DNA (200 PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN (DNA (200 PROTEIN) (DNA ON REPAIR PROTEIN PROTEIN PROTEIN) (DNA ON REPAIR PROTEIN P	DNA-DEPENDENT PROTEIN KINASE (DNA-PK) + DNA DEPENDENT	U35835; [U47077]		-
TELANGIECTASIA (ATM)	PHOLEIN NIVASE CATACTTIC SUBUNIT (DIASTANCS) (ANOCC)		C1a	2250-2680
Color		U33841	C1b	8938-9135
PERDIENT DNA HELICASE II, 70 KD SUBUNIT (LUPLUS KU MAZ865; [S38729] MAZ865; [S38729] MAZ865; [S38729] MAZ865; [S38729] MAZ865; [S38729] MAZ865; [S38729] MAZ865; [S38729] MAZ867; [S70 KD SUBUNIT] CFC MANTIGEN (MUZN) (M		L34075	C1c	6750-7088
NTIGEN PROTEIN P70) (70 KD SUBUNIT OF KU ANTIGEN) NITIGEN PROTEIN P70) (70 KD SUBUNIT OF ED X BINDING PENDENT PURS AUTO-ANTIGEN) DID-LUPUS AUTO-ANTIGEN (TAA) (KUYO) (CTC BOX BINDING PENDENT DID HELICASE II, 86 KD SUBUNIT (LUPUS KU NTIGEN PROTEIN P80) (86 KD SUBUNIT (LUPUS KU NTIGEN PROTEIN P80) (86 KD SUBUNIT) (CTCB) (VUCLEAR FACTOR IV) NTIGEN PROTEIN P80) (66 KD SUBUNIT) (CTCB) (VUCLEAR FACTOR IV) (KRCCS) CASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) GASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) SASA ALPHA GASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) CALIBRAIN PROTEIN RADSO DIJ 13804 C11 DIVIDIA GASE II ALPHA ISOZYME DIJ 13804 C11 DI 12134 DI 12134 C11 DI 12134 DI 12134 C11 DI 12134 C11 DI 12134 C11 DI 12134 C11 DI 12134 DI 12134 C11 DI 12134 C11 DI 12134 DI 12134 C11 DI 12134 DI 12134 C11 DI 12134 C11 D		M32865; [S38729]		
PARTICLE PARTICLE	AUTOANTIGEN PROTEIN P70) (70 KD SUBUNIT OF KU ANTIGEN)			
Pack Pack	(THYROID-LUPUS AUTO-ANTIGEN) (TLAA) (KU70) (CTC BOX BINDING			
PERDIENT DNA HELICASE II, 86 KD SUBUNIT (LUPUS KU M30938) WITGEN PROTEIN PROJEIN PROJEIN PROJEIN PROJEIN PROJEIN PROJEIN PROTEIN PROJEIN CLEAR FACTOR IV) WITGEN PROTEIN PROJEIN (TLAA) (CTC BOX BUINITO C KU ANTIGEN) WAS KD SUBUNIT) (CTCBF)			CIG	1729-1974
INTIGEN PROTEIN P88) (88 KD SUBUNIT OF KU AN IGEN) NITIGEN PROTEIN P89) (88 KD SUBUNIT OF KU AN IGEN) NITIGEN PROTEIN P80) (TCAS) (NUCLEAR FACTOR IV) (XRCCS) (XR		M30938		
Color Colo	AUTOANTIGEN PROTEIN P86) (86 KD SUBUNIT OF KU ANTIGEN) CHYBOID, I IIDI IS ALITOANTIGEN) (TI AA) (CTC BOX BINDING			
CASE CASE	FACTOR RS KID SUBLINITY (CTCBF) (CTCBF) (NUCLEAR FACTOR IV)			
GASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X84740 C1f GASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X84740 C1g GASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X83441 C1h GASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X83441 C1h GASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X83441 C1h GASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X06745 C1i EPAIR PROTEIN RAD50 C1j C1j EPAIR PROTEIN RAD51 HOMOLOG (Replication protein A (E coil D13804 C1j C1j EPAIR PROTEIN RAD52 HOMOLOG U12134 C1j C1j EPAIR PROTEIN RAD52 HOMOLOG U12134 C1j C1j DPOISOMERASE II ALPHA ISOZYME J04088 C1m J04088 NTOSUM GROUP B COMPLEMENTING PROTEIN) (DNA J04088 C1n C2a EPAIR PROTEIN COMPLEMENTING XP-0 CELLS (XERODERMA X5221; [HT1175] C2a C2b EPAIR PROTEIN COMPLEMENTING XP-6 CELLS (XERODERMA IL20046; [X69978]) C2c C2c EPAIR PROTEIN COMPLEMENTING XP-6 CELLS (XERODERMA IL20046; [X69978]) C2c C2c	(KU80) (XRCC5)	,	C1e	2340-2764
GASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X84740 C1g C1g C1g C1h	DNA EXCISION REPAIR PROTEIN ERCC1	M13194	C1f	625-938
C19		X84740		
GASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) X83441 C1h DLYMERASE ALPHA X06745 C1i DLYMERASE ALPHA U63139 C1i EPAIR PROTEIN RAD50 C1i C1i EPAIR PROTEIN RAD52 HOMOLOG (Replication protein A (E coli D13804) C1i EPAIR PROTEIN RAD52 HOMOLOG U12134 C1i DFOISOMERASE I J03250 C1m DFOISOMERASE I JD4088 C1n DFOISOMERASE II ALPHA ISOZYME C1n EPAIR PROTEIN COMPLEMENTING XP-B CELLS (XERODERMA M31899 C1n ON REPAIR PROTEIN COMPLEMENTING XP-D CELLS (XERODERMA X52221; [HT1175] C2a EPAIR PROTEIN COMPLEMENTING PROTEIN) (DNA GOON REPAIR PROTEIN SHOOTEIN) (DNA GOON REPAIR PROTEIN SHOOTEIN) (DNA GOON REPAIR PROTEIN COMPLEMENTING XP-G CELLS (XERODERMA L20046; [X69378]) C2c EPAIR PROTEIN COMPLEMENTING XP-G CELLS (XERODERMA L20046; [X69378]) C2c	(DNL3)		Cig	2460-2780
X06745 C11 U63139 C1 U63139 C1 U12134 C2 U120368 C2 U120368 C2 U20368 U20368 C2 U20368 U20368 C2 U20368 U20368 C2 U20368 U203	DNA LIGASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP))	X83441	-	
No6745 C1i	(DNL4)		C1h	2787-3074
Replication protein A (E coli D13804 C1j Replication protein A (E coli D13804 C1k U12134 C1i U03250 C1m J04088 C1n P-B CELLS (XERODERMA M31899 C1n G PROTEIN) (DNA C2a P-D CELLS (XERODERMA X52221; [HT1175] C2a G PROTEIN) (DNA C2b M36089 C2c G PROTEIN) (DNA C2c G PROTEIN) (DNA C2c	DNA POLYMERASE ALPHA	X06745	C1i	3721-4093
Replication protein A (E coli D13804 C1K C1K C1 C2 C3	DNA REPAIR PROTEIN RADSO	U63139	C1j	5117-5435
P-B CELLS (XERODERMA M31899 G PROTEIN) (DNA H 89 KD SUBUNIT) G PROTEIN) (DNA C2a P-D CELLS (XERODERMA X52221; [HT1175] G PROTEIN) (DNA M36089 C2b M36089 G2b G2b GP-G CELLS (XERODERMA L20046; [X69978]	DNA REPAIR PROTEIN RAD51 HOMOLOG [Replication protein A (E coli	D13804		
U12134 C11 U3250 C1m U30850 C1n U3088 C1n U3088 C1n P-B CELLS (XERODERMA M31899 C2a P-D CELLS (XERODERMA X5221; [HT1175] G PROTEIN) (DNA C2b M36089 C2c C2b C2b C2b C2b C2c C3c C4c	RecA homolog, RAD51 homolog)	•	Cik	867-1159
J03250 C1m J04088 C1n J04088 C1n J04088 C1n L RANSCRIPTION H 89 KD SUBUNIT) C2a P-D CELLS (XERODERMA X52221; [HT1175] C2b G PROTEIN) (DNA C2c C2b C2c C2b C2c C2b C2c C2b C2c C2b C2c C2c C2c C2c C2c C2c C2c C2c C2c C2c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c C3c	DNA REPAIR PROTEIN RAD52 HOMOLOG	U12134	C1	1528-1733
P-B CELLS (XERODERMA M31899 G PROTEIN) (DNA L TRANSCRIPTION H 89 KD SUBUNIT) C2a P-D CELLS (XERODERMA X52221; [HT1175] G PROTEIN) (DNA M36089 C2b M36089 C2c P-G CELLS (XERODERMA L20046; [X69978]	DNA TOPOISOMERASE I	J03250	C1m	2388-2796
C2a [HT1175] C2b C2b [X69978]	DNA TOPOISOMERASE II ALPHA ISOZYME	J04088	Ofn	2459-2883
C2a C2b C2c	DNA-REPAIR PROTEIN COMPLEMENTING XP-B CELLS (XERODERMA	M31899		
C2a C2b C2c	PIGMENTOSUM GROUP B COMPLEMENTING PROTEIN) (DNA			
C2a C2b C2c	EXCISION REPAIR PROTEIN ERCC3) (BASAL THANSCHIPTION			
C2b C2c	FACTOR 2 89 KD SUBONIT) (BIFZ-p89) (Trim 89 KD SUBONIT)		C2a	2109-2466
MACC-2 C2b MAG089 C2c	DNA-REPAIR PROTEIN COMPLEMENTING XP-D CELLS (XERODERMA	X52221; [HT1175]		
EMENTING XP-G CELLS (XERODERMA L20046; [X69978]	PIGMENTOSUM GROUP D COMPLEMENTING PHOLEIN) (DNA		-	1001
EMENTING XP-G CELLS (XERODERMA L20046; [X69978] WPLEMENTING PROTEIN) (DNA	EXCISION REPAIR PROTEIN ERCC-2)		CZD	1220-1821
Ţ	DNA-REPAIR PROTEIN XRCC1	M36089	C2c	1226-1539
750	DNA-REPAIR PROTEIN COMPLEMENTING XP-G CELLS (XERODERMA	L20046; [X69978]		
ככם	FIGURE NECESTARY OF COURT EXPERIENCE COURTS (COURT) (COURT)		C2d	1374-1638

TABLE 3 (CONT)

		A	Doctition
	Gendank #	Allay cooluliate	100000
GE-INDUCIBLE PROTEIN E PROTEIN (CHOP).	S40706 [S62138]	C2e	480-789
GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD45 M60974 IDNA-DAMAGE INDUCIBLE TRANSCRIPT 1) (DDIT1).	M60974	C2f	526-886
RANSFERASE (6-	M29971	C2g	241-546
	X90392; [L40817; U06846]	CZh	2038-2427
DAMA MICHATCH BEDAIR PROTEIN MI H1 (mint HOMOLOG)	U07418	CZi	1765-2020
	124564	C2j	489-780
VATOR 1 36 KD SUBUNIT (REPLICATION FACTOR C 36 KD IMP (REC36)	L07540	C2k	708-1051
KD SUBUNIT (REPLICATION FACTOR C 37 KD	M87339	C2I	98-355
KD SUBUNIT (REPLICATION FACTOR C 38 KD	L07541	C2m	438-762
KD SUBUNIT (REPLICATION FACTOR C 40 KD	M87338	C2n	882-1286
TEIN A 70 KD DNA-BINDING SUBUNIT (RP-A) (RF- ACTOR-A PROTEIN 1) (SINGLE STRANDED DNA-	M63488		1498-1838
Sinconvince Distall (Superoxide dismutase 1 (Cu/Zn)	HT3218 [K00065]	C3P	198-496
	M96684	C3c	563-855
HHRBA (YEAST RADS HOMOLOG) (UBIQITIN-CONJUGATING ENZYME) (UBCA)	M74524	C3d	175-433
UV EXCÍSION REPAIR PROTEIN PROTEIN RAD23 [xeroderna pigmentosum group C repair complementing protein HHR23A]	D21235	C3e	355-632
CELL FATE/DEVELOPMENT REGULATORS			
-Notch pathway	M73980	C3(2701-2965
Notes	U77493	C3g	373-658
potch around protein (N)	M99437	C3h	647-1210
North A	U95299	C3i	3014-3169
Langed 1	AF028593	C3j	3884-4117
Jacobed 2	AF003521	C3k	1027-1241
DELTA-LIKE PROTEIN PRECURSOR (CONTAINS: FETAL ANTIGEN 1) (FA1) (DLK) + ADRENAL SPECIFIC 30Kd PROTEIN GB: X17544	U15979; [Z12172]	C3I	1090-1403
manic fringe	U94352	C3m	979-1235

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
lunatic fringe	U94354	C3n	563-857
-Wnt pathway			
WNT2 OR IRP	X07876	C4a	899-1252
Wnt-5a	L20861	C4b	1036-1281
WNT-8B	X91940	C4c	164-447
WNT-108	X97057	C4d	330-635
Wni-13	271621	C46	569-847
frizzled	L37882	C4f	1491-1756
inzzled-related FrzB (Fritz) (frezzled (fre))	U24163; [U91903; U68057]	C4g	590-819
frizzled 5	U43318	C4h	936-1091
(FZD3)	U82169	C4i	865-1182
dishevelled (DVL) + dishevelled 3 (DVL3)	U49262; [U75651]	C4j	1311-1610
dishevelled homolog (DVL)	U46461	C4k	1409-1586
-Hedgehog pathway		,	
sonic hedgehog (SHH)	L38518	C41	164-474
patched homolog (PTC)	U43148	C4m	3179-4050
smoothened	U84401	C4n	503-789
RECEPTORS			
5T4 ONCOFETAL ANTIGEN	Z29083	CSa	748-981
AXL (TYROSINE-PROTEIN KINASE RECEPTOR UFO)	M76125	CSb	2045-2348
GATION-INDEPENDENT MANNOSE-6-PHOSPHATE RECEPTOR lines like growth factor resentor II. IGFR-21	Y00285; [J03528]	C5c	1394-1831
CDW40; NERVE GROWTH FACTOR RECEPTOR-RELATED B-	X60592	550	198-605
EMPHOCT IE ACHIVALION MOLECOLE FPIDERMAI GROWTH FACTOR RECEPTOR PRECURSOR (EC	K03193; [X00588; X00663;		
2.7.1.12). (EGFR) (ERBB1)	U48722]	CSe	3410-3757
EPS 15 (AF-1P PROTEIN)	U07707; [Z29064]	CSf	1828-2140
EPS8	U12535	C5g	2293-2645
ERBB4	L07868	CSh	3570-3965
ERYTHROPROTEIN RECEPTOR	M60459	CSI	1423-1740
FAU	X65923	CSi	8-344
GARP	724680	C5k	3399-3777
HER2 (ERB-B2)	M11730; [M95667]	CSI	2556-2722
HER3 (ERB-83)	M29366; [M34309]	C5m	3886-4139
HGF ACTIVATOR	D14012	CSn	1487-1845
HGF ACTIVATOR LIKE	D49742; [S83182]	C6a	311-595
IGFBP COMPLEX ACID LABILE CHAIN	D25216	C6b	1509-2669
IGFBP2	M35410	066	680-1071

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
IGFBP3 (GROWTH HORMONE-DEPENDENT INSULIN-LIKE GROWTH EACTOR RINDING PROTFIN)	M31159; [M35878]	C6d	451-744
IGEBP4	M62403	C6e	657-967
(GEBP5	M65062	Cef	356-602
IGEBP6	M62402	Ceg	345-536
INSUITING INCOME BECTOR I RECEPTOR	X04434	Ceh	3413-3904
BASIC FIBROBLAST GROWTH FACTOR RECEPTOR 1 PRECURSOR	M37722; (X66945;		
(BFGF-R) (EC 2.7.1.112) (FMS-LIKE TYROSINE KINASE-2) (C-FGR) (FGFR) (FIG) (FGFR) (FLT2). (HBGF-R-ALPHA-A1) (HBGF-R-ALPHA-	M63887; M63888; M63889;M34186; M34641]		
A2) (HBGF-R-ALPHA-A3) + FGFR SECRETED FORM (M34188)			1746-1967
NEBVE GROWTH FACTOR RECEPTOR	M14764	Cej	2762-3242
PNGER-AI PHA	M21574	C6k	5118-5583
POGER-BETA	M21616	Cel	842-1133
transmembrane receptor precursor (PTK7); COLON CARCINOMA KINASE-A (CCKA)	U33635; [U40271]	Cem	3507-3784
SEX GENE	X87852	Cen	209-433
TRANSFORMING GROWTH FACTOR-BETA TYPE III RECEPTOR	L07594	C7a	3358-3592
THANSMEMBRANE PROTEIN TMP21	X97442	C7b	380-1176
HIGH AFFINITY NERVE GROWTH FACTOR RECEPTOR PRECURSOR	X03541		
(EC 2.7.1.112) (TRK1 TRANSFORMING TYROSINE KINASE PROTEIN)			
(P140-1HKA) + tR-13 (PBB 1HK-13 ONCOPROTEIN)		C7c	1816-2118
Irk-T3 (P68 TBK-T3 ONCOPROTEIN)	X85960	C7d	252-1112
Irrk-B	U12140	C7e	1006-1384
I-K-C	U05012	C7f	359-765
TUMOR NECROSIS FACTOR RECEPTOR 1	M33294	C7g	1570-1817
TUMOR NECROSIS FACTOR RECEPTOR 2 PRECURSOR (TUMOR	M32315; [M55994]		
NECROSIS FACTOR BINDING PROTEIN 2) (TBPII) (P80) (TNF-R2) [P75) (CD1208) (TNFB2) (TNFBR).		C7h	3359-3543
RETINOIC ACID RECEPTOR ALPHA1 (RAR- ALPHA1) + PML-RAR	M73779; [X06538; [X06614]	C7i	2935-3238
retiroic acid receptor alpha [RETINOIC ACID RECEPTOR RXR-ALPHA (PERPA)	X52773	C7i	352-616
retinoic acid receptor epsilon [RETINOIC ACID RECEPTOR BETA-2 (RAR X07282; [Y00291] BETA-2) (RAR-EPSILON)	X07282; [Y00291]	C7k	1315-1633
retinoic acid receptor gamma [RETINOIC ACID RECEPTOR GAMMA]	M24857; [M38258; M57707; M32074]	C71	1569-1834
retinoic acid receptor rxr-beta [RETINOIC ACID RECEPTOR RXR-BETA]	M84820; [X63522]	C7m	643-1135

TABLE 3 (CONT)

		C7n 5117-5435	5117-5435
THROMBOPOELLIN RECEPTION			
QUADRANT D			
CELL ADHESION MOTILITY, AND INVASION			
RE PROTEIN (CSPCP)	M55172		
(AGGRECAN 1)(CHONDROITIN SULFATE PROTEOGLYCAN CORE		018	6705-6956
(17	104599	D1b	854-1129
an		D1c	296-960
		D1d	105-1163
CHONDROITINDERMATAN SULFATE PROTEOGLYCAN CORE	M14219	à	71.2 806
PROTEIN (DECORIN) (PG-S2) (PG40)		Ule	060-717
COLLAGEN (LEGONRP)	D21337	D1f	5342-5588
COLEMEN TOOLS	X55525	D1g	428-741
	X16468	D1h	3604-3751
Collagell type if alpha 1	X14420	D1i	3867-4046
collagen type III pro-alpha-1		D1j	882-1113
Collagen type ty alpha a		D1k	2296-2545
congentype to apria-3	X15879	D1I	316-688
collagen type vi apria-1	M34570	D1m	203-396
collagell type vi aprime c		D1n	640-1487
Collabell type VI alpha 1	X57527	D2a	612-1772
	J04177	D2b	2864-3091
	U32169	D2c	4473-4769
Collagen type At pro-alpha-2	M92642	D2d	4816-5991
collaboration and a series of the series of	L22548	D2e	2300-2539
LALDAN ANGAN	X70904; [X91171]	D2f	1018-1388
LAMON (CAMA)	S77512	D2g	3871-4158
LAMBZ (CAMILLIN)	M61916	DZh	3177-3554
Idrimini Di	J03202	D2i	2878-3232
laminin BC	U43901	D2j	460-812
יייייייייייייייייייייייייייייייייייייי	U86759	D2k	859-1147
The state of the s	M30269	D2I	2120-2428
TOURSOIN	X78565	D2m	6652-6924
TENANCIA: O	X98085	D2n	3916-4165
VERSICAN [isoforms , V1, V2, V3]	U16306; [X15998; U26555;		100 074
	D32039	D3a	1103-374

TABLE 3 (CONT)

			Bocition
Cell Cycle/Growth Regulators	Genbank #	Array Coolumbate rosition	TOSING L
SPARC PRECURSOR (SECRETED PROTEIN ACIDIC AND RICH IN	J03040		
CYSTEINE) (USTEUNEUTIN) (UN) (BASEMEINT MEMBERSETTET)		D3b	280-642
TUDONIDOCEDANDIN 1 PRECIESOR	X14787	D3c	3187-3450
TUDOMBOSPONDIN 9 PRECIBEOR	L12350	D3d	3151-3531
VITRONECTIN PRECURSOR (SERUM SPREADING FACTOR) (S-	X03168		0007
PROTEIN (CONTAINS: SOMATOMEDIN B)		U3e	3/21-4093
(ibroactio	X02761	D3f	6163-7290
DNA binding protein Hel-N2: Ft AV-like neuronal protein 1	U12431; [U29943]	D3g	1006-1384
LEGADAN SHI FATE PROTECISI YCAN (HSPG2)	M85289	D3h	1232-1389
integrals alpha	X68742	D3i	2690-2976
integrin alpha? [very late antigen-2 (vla-2)/collagen receptor alpha-2	M28249; [X17033]	Dai	2367-2664
Subunit	M59911	D3k	2564-2944
Integrin alphas	L12002: [X16983]	D3I	2709-3063
Integrin alpha 4	X06256	D3m	2094-2367
٦	X53586; [X59512]	D3n	3642-3988
imegnii alpiido	X74295	D4a	255-591
Integrin alpha o	L36531	D4b	2709-3063
megini aprida	D25303; [L24158]	D4c	706-980
integrill appliab	L25851	D4d	2279-2529
inform bata 1	M34189	D4e	701-1301
integrin beta 7 PI ATEI FT MEMBRANE GLYCOPROTEIN IIIA]	J02703; [M25108]	D4f	2038-2373
integrity betad	X53587; [X52186]	D4g	5357-5697
integrity Detail	J05633	D4h	2279-2528
intendin beta6	M35198	D4i	1619-1901
integrin beta7	M62880	D4 i	2562-2944
integrin beta8	M73780	D4k	22-877
Foral adhesion kinase	L13616	041	2179-2631
Integrin-linked kinase (ILK)	U40282	D4m	1245-1530
Protein tyrosine kinase Pyk2 (Cell adhesion kinase-beta, CAK-beta) (FAK2) U43522; [L49207]	U43522; [L49207]	D4n	3658-3952
- 111.	U14588	D5a	1260-1644
raxiiin	X94991: [X95735]	D5b	585-1514
Zyzin + Zyzin-Z	AF000974	D5c	1240-1466
Lyail related protein Lin -	U37139	DSd	606-1504
Oeta Steffworth ike profein	U59752	D5e	43-338
	M38690	DSf	372-962
Ezip (cdoxillin 2)	X51521	DSg	1611-1883

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
MERLIN (SCHWANNOMIN) (moesin-ezrin-radixin-like	L11353; Z22664; X72657;		
protein)(neurofibromatosis 2)	L27133	Osh	355-674
L1CAM	M74387	DSi	3197-3485
N-CAM INEURAL CELL ADHESION MOLECULE,	X16841	:50	2000 0046
PHOSPHA I IDYLINOSI I OL-LINKED ISOPOHM; CUSE		00	2330-2040
NINJURIN-1	U72661	D5k	212-492
opioid binding cell adhesion molecule	L34774	DSI	115-728
DCC	X76132	D5m	893-1189
P37NB	U32907	DSn	95-456
PLEXIN	U52111	D6a	585-1514
semaphorin (CD100)	00800	Deb	2517-2921
semaphorin E	AB000220	Dec	2949-3181
semaphorin (ii	126081	. peq	899-1152
semaphorin V	U33920	D6e	177-442
SEMAPHORIN-1	U38276	Def	488-653
TAX1, AXONIN-1/TAQ1	X85978	Deg	209-433
LAR	Y00815	Deh	5799-6049
HYALURONAN RECEPTOR (RHAMM)	U29343	Dei	2496-2798
PLATELET GLYCOPROTEIN IV (GPIV) (GPIIIB) (CD36 ANTIGEN) (PAS	M24795		
IV) (PAS-4 PROTEIN)		Dej	554-806
caveolin-2	AF035752 U32114	D6k	1340-1519
caveolin-1	Z18951 S49856	Del	62-413
ANGIOGENESIS REGULATORS			
VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR 2	L04947; [X61656]		
PRECURSOR (EC 2.7.1.112) (VEGFR-2) (KDR) (KINASE INSERT			
DOMAIN RECEPTOR) (FRAGMENT)		D6m	2686-3053
VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR 3	X68203; [X69878; U43143]		
PRECURSOR (EC 2.7.1.112) (VEGFR-3) (TYROSINE-PROTEIN KINASE IRECEPTOR EL T4. CLASS III).		Den	4236-4402
FL CYTOKINE RECEPTOR PRECURSOR (EC 2.7.1.112) (TYROSINE-	U02687		
PROTEIN KINASE RECEPTOR FLT3) (STEM CELL TYROSINE KINASE			7000
TYBOSINE-PROTEIN KINASE BECEPTOR TIE-1 PRECURSOR (EC	X60957 [S89716]	חימ	2431-2905
2.7.1.112).		D7b	3114-3536
TYROSINE-PROTEIN KINASE RECEPTOR TIE-2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN KINASE RECEPTOR TEK) (P140 TEK)	L06139		
(TUNICA INTERNA ENDOTHELIAL CELL KINASE).		D7c	3243-3586

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
VASCULAR ENDOTHELIAL GROWTH FACTOR B PRECURSOR (VEGF-1U48801; [U43368] B) + VEGF RELATED FACTOR ISOFORM VRF186 PRECURSOR	U48801; [U43368]		
		D7d	158-648
VASCULAR ENDOTHELIAL GROWTH FACTOR C PRECURSOR (VEGF- U43142 C) (VASCULAR ENDOTHELIAL GROWTH FACTOR RELATED	U43142		
PROTEIN) (VRP) (FLT4 LIGAND).		D7e	1165-1559
PLACENTA GROWTH FACTORS 1 AND 2 PRECURSOR (PLGF-1/	X54936		
PLGF-2).		D7f	1098-1371
SL CYTOKINE PRECURSOR (FLT3/FLK2 LIGAND).	U04806; [U03858]	D7g	29-362
angiopoietin-1	U83508	D7h	1749-2031
CH FIBROBLAST GROWTH FACTOR RECEPTOR [Golgi loakcoprotein MG160 (GLG1)]	U28811; [U64791]	D7i	3279-4140
FGFR3 (FLG-2)	M58051; [X58255]	07	323-896
FGFR4	L03840	D7k	1503-1743
FIBROBLAST GROWTH FACTOR RECEPTOR 2 PRECURSOR (FGFR-	U11814; [M80634; X52832; M35718; M87771; M87772]		
(FGFR2) (BEK) (BFR-1) (KSAM-1) + K-SAM; K-SAM-III; K-SAM-IV		IZO	753-1189
VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR 1	U01134; [X51602]		
PRECURSON (EC Z./1.112) (VEGFR-1) (1780SINE-FRO LEIN NINASE RECEPTOR FLT) (FLT-1) (SFLT)		D7m	1288-1604
HOMEOBOX PROTEIN HOX-D3 [HOX 4A]	D11117	D7n	4200-4447
QUADRANTE	-		
INVASION REGULATORS			
MMP-1 (collagenase-1)	X05231	E1a	512-836
MMP-2 (gelatinase A)	J03210, [J05471]	E1b	477-778
MMP-3 (stromelysin-1)	X05232	E1c	331-1491
MMP-7 (matrilysin)	X07819	E1d	335-738
MMP-8 (collagenase-2)	J05556	E1e	532-865
MMP-9 (gelatinase B)		E1f	1012-1346
MMP-10 (stromelysin-2)	[M30461]	E1g	387-1319
MMP-11 (stromelysin-3)		E1h	263-1508
MMP-12 (metalloelastase)	L23808	E1i	275-787
MMP-13 (collagenase-3)	X75308	E1j	463-761
MMP-14 (MT1-MMP)	D26512, [X83535]	Eik	413-749
MMP-15 (MT2-MMP)	248482	==	1210-1456
MMP-16 (MT3-MMP)	D50477	E13	991-1226
MMP-17 (MT4-MMP)	X89576	E1n	630-1830
MMP-19	X92521	E2a	1383-1655

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate Position	Position
EPA)	X03124	E2b	194-492
	105593	E2c	403-694
inducible gane 5 mig-5)	230183	E2d	346-587
	U76456	E2e	445-671
overscellular matrix metalloproteinase inducer EMMPRIN	120471	E2f	23-354
UROKINASE-TYPE PLASMINOGEN ACTIVATOR PRECURSOR (EC	M15476	E20	824-1120
13.4.21.73 (UPA) (U-PLASMINOGEN ACTIVATOR PRECURSOR (EC	M15518; [X07393;	6	
13 4 24 68) T-PA (T-PI ASMINOGEN ACTIVATOR).	M18182]	Ezh	1221-1577
DI ASMINOGEN PRECIPSOR (EC 3.4.21.7)	X05199	E2i	1859-2162
PLASMINOGEN ACTIVATOR INHIBITOR-1 PRECURSOR,	X04429	icu icu	1105.1242
ENDOTHELIAL (PAI-1)		[27]	10010
PLASMINOGEN ACTIVATOR INHIBITOR-2, PLACENTAL (PAI-2)	M18082;[J02685]	E2k	378-954
PLASMA SERINE PROTEASE INHIBITOR PRECURSOR (PCI)	M68516; [J02639]		
(PROTEIN C INHIBITOR) (PLASMINOGEN ACTIVATOR INHIBITOR-3)		č	0000 0000
(PAI3).		EZ	0740-0000
UROKINASE PLASMINOGEN ACTIVATOR SURFACE RECEPTOR, GPI-	U08839 [M83246; X51675]		
ANCHORED FORM PRECURSOR (U-PAR) (MONOCY I E ACTIVATION		F2m	749-1043
ANTIGEN MO3) (CD8/ ANTIGEN)	0,000		
LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PHOLEIN 1 PRECURSOR (LRP) (ALPHA-2-MACROGLOBULIN RECEPTOR) (A2MR)	91851X		
		E2n	5439-5742
LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 2	U04441	п 33	1365-2162
(MEGALIN) (GLYCOPHOLEIN 330) (PRAGMENT)	M111313	Eah	3972-4325
ALPHA-Z-MACHOGLOBOLIN PRECURSON (ALTIAZZ-W) DI ATELET BASIC PROTEIN PRECURSON (PBP) (CONTAINS:	M54995; M38441		
CONNECTIVE-TISSUE ACTIVATING PEPTIDE III (CTAP-III), LOW-			
AFFINITY PLATELET FACTOR IV (LA-PF4), BETA-			,
THROMBOGLOBULIN (BETA-TG), NEUTROPHIL-ACTIVATING	***		
PEPTIDE 2 (NAP-2))		E3c	63-252
ALPHA-2-MACROGLOBULIN RECEPTOR-ASSOCIATED PROTEIN	M63959		
PRECURSOR (ALPHA-2-MHAP) (LOW DENSITI LIPOTHOLICINA)		E3d	440-890
HECEP ION-RELATED FROILIN-ASSOCIATED FROIL 1 / 1 / 1 / 1 / 1 / 1 / 1 / 1 / 1 / 1	X17620		
KINASE A) (TUMOR METASTATIC PROCESS-ASSOCIATED PROTEIN)			
(METASTASIS INHIBITION FACTOR NM23) (NM23-H1).		E3e	245-612

TABLE 3 (CONT)

	Gon Rank #	Array Coordinate Position	Position
	L16785; [M36981]		
KINASE B) (NM23-H2) (C-MYC PURINE-BINDING TRANSCHIPTION		E3f	69-351
LEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6)	Y07604	E30	141-448
1000	U43527	E3h	116-454
malignant melanoma melasiasis-suppressor (NOC-1) gono	U35113	E3i	957-1825
F ANTIGEN (PSM)	M99487	E3j	1068-1200
integrin/cysteine-rich protein precursor (MDC9)	U41766	E3k	640-958
	X06820	E3I	53-1648
tho (Ha) SMALL GTPase (rhoC).	L25081		637-1473
. 1	X61587		900-1228
Don't de la contraction de la	Y07923	E4a	33-388
	X95456	E4b	75-377
nivor protein	X95282	E4c	209-534
ED C3 BOTULINUM TOXIN SUBSTRATE 1 (P21-RAC1)	M29870; [M31467]	77	EE. 420
(RAS-LIKE PROTEIN TC25)		T to	2014
RAS-RELATED C3 BOTULINUM TOXIN SUBSTRATE 1 (P21-RAC2)	M64595; [M29871]	E48	31-1100
ras-like protein TC10	M31470	E41	005-02
res-like small GTP ase TTF	Z35227	E4g	491-759
HOTHER CONTRACTOR	D85815	E4h	130-361
Des assessing a collection containing profess presse pressed the profession of the pressed of th	U43195	E4i	3793-4233
COCAS CTDsco-artivation protein	U02570	E4j	864-1182
Col disconstitution RhoGD loamma	U82532	E4k	309-554
Thurstone invesion and metastasis inducing TIAM1	U16296	E41	4275-4645
PUTATIVE RHO/RAC GUANINE NUCLEOTIDE EXCHANGE	U11690	Į.	3033 4165
FACTOR(RHO/RAC GEF) (FACIOGENITAL DYSPLASIA PROTEIN)	1,70017	E4m	781-1170
RHO-GAP HEMATOPOIETIC PROTEIN C1 (P115) (KIAA0131).	1,0000	100	322.600
rho GDP-dissociation inhibitor protein 2 (Ly-GDI)	L20688	ESa	226-200
πο GDP-dissociation Inhibitor 1	X69550	CSD	328-024
SERINE/THREONINE-PROTEIN KINASE PAK-ALPHA (EC 2.7.1) (P65-	U24152	ii C	756-1055
PAK) (P21- ACTIVALED KINASE) (ALTHA-TAN)	1124153	E5d	335-671
p21-activated protein kinase (Pakz)	2015		
CADHERIN-2 (N-CADHERIN)	M34064 [X57548; X54315]	E5e	942-1299
	0.5550	100	542.R35

TABLE 3 (CONT)

	Condon't	Array Coordinate Position	Position
Cycle/Growth Hegulators IERIN-4 RETINAL-CADHERIN PRECURSOR (R-CADHERIN) (R-	L34059	E5a	1172-1425
CADHERIN-5 VASCULAR ENDOTHELIAL-CADHERIN PRECURSOR (VE-X79981; [X59796]	X79981; [X59796]	E5h	1607-1769
CADHERIN) (784 AN IGEN) (CD144 AN IGEN).	031784	ESi	2119-2443
	34060	ESi	1069-1347
	134056	E5k	1778-2076
CADHERIN 12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL	L34057; [L33477]	E5I	657-903
CADHERIN-13 T-CADHERIN PRECURSOR (TRUNCATED-CADHERIN)	L34058; [U59289; U59288]	E5m	949-1187
CADHERIN-14 MUSCLETTIN PRECURSOR (M-CADHERIN)	D83542	Esn	228-456
(CADHERIN-14) (CADHERIN-13) ALPHA-CATENIN (CADHERIN-ASSOCIATED PROTEIN) (ALPHA E-	D13866 [D14705 L23805;	E6a	55-492
CATENIN)	M94151	E6b	2296-2545
ALPHA-CATENIN HELATED PROTEIN (CATENIN ALTER)	X87838 [219054]	E6c	2061-2463
BETA-CATENIN	M23410	E6d	2000-2312
PLAKOGLOBIN (DESMOTLANIN III)	M74088; [M73548]	E6e	7992-8326
neuroendocrine-dig (NE-dig) a novel human homolog of the Drosophila neuroendocrine-dig (NE-dig) a novel human homolog with the APC protein	U49089		
discs large (dlg) tumor suppressor protein interacting with the 2th of process		E6f	2210-3116
ORA - Transfer in the ABC	U24166	E6g	488-796
EB1, a protein that binds to APC	L11370	E6h	1246-1605
protocadhenn 42	L11373	E6i	1018-1388
protocadherin 43	M77830	E6j	6987-7826
desmoplakin I	U53786	E6k	5583-5788
envoplakin (EVPL)	M63618	E6l	5680-6055
bullous pempingola antigen	Z26317 [S64273]	E6m	2819-3135
desmoglein z	X56654	E6n	2578-2889
desmogrent type :	X72925	E7a	475-1154
desmocollin type 1	X83929; [D17427]	E7b	608-1607
desmocollin type 3 + destructural type 4	X56807	E7c	802-1115
EPHRIN-A1 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE	M57730 M37476		
KINASE LIGAND 1) (LERK-1) (IMMEDIATE EARLY RESPONSE			
PROTEIN B61) (TUMOR NECROSIS FACTOR, ALPHA-INDUCED		E7d	124-1062
EPHRIN AS PRECURSOR (EPH-RELATED RECEPTOR TYROSINE	U26403	E79	375-1325
KINASE LIGAND 7) (LEMN-7) (AL-1).			

TABLE 3 (CONT)

	4 7 2 0	Arran Coordinate	Docition
Cell Cycle/Growth Regulators	Genbank #	Allay Cooldinate rosition	LOBIGOL
EPHRIN-BI PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE I IGAND 2) (I ERK-2) (ELK-LIGAND PRECURSOR) (ELK-LI).	U09304	E7f	507-1186
EPHRING-BY PRECINE (FPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 5/11 FRK-5) (HTK LIGAND) (HTK-L).	L38734	E79	442-560
EPHRIN-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) (LERK-8) (EPH-RELATED RECEPTOR	U66406	E7h	2056-2282
EPHRIN TYPE-A RECEPTOR 2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN KINASE RECEPTOR ECK) (EPITHELIAL CELL	M59371 M36395	Ï	240 4426
KINASE). EPHRIN TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN KINASE RECEPTOR EHK-1) (EPH HOMOLOGY	X95425	[C/1	644-1300
KINASE-1) (RECEPTOR PHOTEIN- LYHOSINE KINASE HENZ). EPHRIN TYPE-B RECEPTOR 1 PRECURSOR (EC 2.7.1.112)	L40636	E7k	998-1469
EPHRIOINETTION ENTRY PRECURSOR (EC 2.7.1.12) TYPE-BROTEPL-BROTE THE CURSOR (EC 2.7.1.12) TYPE-BROTEPL-3 (DRT)	L41939	E71	454-1225
EPHRIN TYPE-B RECEPTOR 4 PRECURSOR (EC 2.7.1.112) TYPE-BROTEIN KINASF RECEPTOR HTK).	U07695	E7m	756-1652
TYROSINE-PROTEIN KINASE HCK (EC 2.7.1.112) (P59-HCK AND P60-HCK) (HEMOPOIETIC CELL KINASE).	M16591	E7n	194-1187
QUADRANT F			
GROWTH FACTORS/CYTOKINES		i	744 007
AMPHIREGULIN	M30704	F12	43-248
BCGF1 (B-cell growth factor)	M61176	F1c	982-1265
DOINT RETA NGE	X52599	F1d	360-1339
VASCULAR ENDOTHELIAL GROWTH FACTOR PRECURSOR (VEGF) (VASCULAR PERMEABILITY FACTOR) (VPF).	M32977; [M27281]	F1e	198-622
BIGH3	M77349	F11	705-1703
BONE MORPHOGENETIC PROTEIN 1 (procollagen C-proteinase) (pCP- M22488; [U50330]	M22488; [U50330]	F19	702-1098
PONE MORPHOGENETIC PROTEIN 2A	M22489	F1h	567-997
BONE MORPHOGENETIC PROTEIN 3	M22491	F1i	1458-1731
BONE MORPHOGENETIC PROTEIN 3B	D49493	F1j	16188-16418
BONE MORPHOGENETIC PROTEIN 4 (BMP-2B)	[D30751; [M22490]	F1k	943-1321
BONE MORPHOGENETIC PROTEIN 5	M60314	F11	1679-1982

TABLE 3 (CONT)

Cell Cycle/Growth Regulators	GenBank #	Array Coordinate	Position
RONF MORPHOGENETIC PROTEIN 6	M60315	F1m	1067-1327
BONE MORPHOGENETIC PROTEIN 7 (OSTEOGENIC PROTEIN 1)	M60316	F1n	451-691
RONE MORPHOGENETIC PROTEIN 8 (OSTEOGENIC PROTEIN 2)	M97016	F2a	1345-1645
BPGE.1	L42379	F2b	825-1213
CNTF. ISOFORM B AND C	A26792	F2c	213-448
CONNECTIVE TISSUE GROWTH FACTOR	M92934	F2d	1459-1748
FGF (kidney)	X04571	F2e	4164-4434
FGE-LIKE GROWTH FACTOR	M60278	F2f	1905-2146
andothelin 2	M65199	F2g	338-570
andothelin 3	J05081	F2h	1428-1685
HEPARIN-BINDING GROWTH FACTOR 1 PRECURSOR (HBGF-1)	X51943; [M13361; X65778]		
(ACIDIC FIBROBLAST GROWTH FACTOR) (AFGF) (BETA-		č	7,00
ENDOTHELIAL CELL GROWTH FACTOR) (ECGF- BETA).		F21	7001-1611
FGF2; HEPARIN-BINDING GROWTH FACTOR 2 PRECURSOR	M27968		
(PROSTATROPIN): (Tage 2) (BASIC TIETTOBENS) (STEERS) (PROSTATROPIN)		F2j	1384-1646
FGF-3: INT-2 PROTO-ONCOGENE PROTEIN PRECURSOR	X14445		
(FIBROBLAST GROWTH FACTOR-3)(HBGF-3).		F2k	189-940
FGF-5: FIBROBLAST GROWTH FACTOR-5 PRECURSOR (HBGF-5).	M37825	F2I	603-1086
FGF-6, FIBROBLAST GROWTH FACTOR-6 PRECURSOR (HBGF-6)	X63454	1	207 466
(HST-2).		rzm	004-707
FGF-7; KERATINOCYTE GROWTH FACTOR PRECURSOR (KGF) (FIBRORI AST GROWTH FACTOR- 7) (HBGF-7).	M60828	F2n	522-955
FGF-8, ANDROGEN-INDUCED GROWTH FACTOR PRECURSOR	U36223	,	000
(AIGF) (HBGF-8) (FIBROBLAST GROWTH FACTOR-8)		⊩3a	32-3100
FGF-9, GLIA-ACTIVATING FACTOR PRECURSOR (GAF) (FIBROBLAST	D14838	F34	110.949
GROWIN FACTOR-9) (nBGP-9).	1166197	F3c	17-566
TONE	L19063	F3d	248-390
GLIA MATURATION FACTOR beta	HG563 [M86492;		
	AB001106]	F3e	203-434
RECOMBINANT GLIAL GROWTH FACTOR + NEU DIFFERENTIATION	L12260; U02326; M94165	č	4000
FACTOR + HEREGULIN		rsi	1009-1407
TRANSFORMING GROWTH FACTOR-BETA-2 (glioblastoma-derived total suppressor factor)	M19154; [Y00083]	F3g	1538-1878
GROWTH INHIBITORY FACTOR (METALLOTHIONEIN-III) (MT-III)	D13365; [M93311]	F3h	4-1052

TABLE 3 (CONT)

PARIN-BINDING GROWTH- PARIN-BINDING GROWTH- CIFIC FACTOR 1) (OSF-1) TH PROMOTING FACTOR 1) I (EGR-1) (KROX24) IC FINGER PROTEIN 225) IC FINGER PROTEIN 225) IC Macrophage-stimulating	M57399; [X52946; D90226]		
÷	57399; [X52946; D90226] [62829; [X52541]		
£	162829; [XS2541]		
÷	(62829; [XS2541]		
£ .	162829; [XS2541]		
I (EGR-1) (KROX24) IC FINGER PROTEIN 225) (macrophage-stimulating	l62829; [XS2541]		
i (EGR-1) (KROX24) IC FINGER PROTEIN 225) (macrophage-stimulating	162829; [X52541]	F3i	602-847
IC FINGER PROTEIN 225) (macrophage-stimulating			
(macrophage-stimulating			
(macrophage-stimulating		F3j	989-1276
	174178		
proper (MST1))		F3k	1643-2015
HEPTOMA-DERIVED GROWTH FACTOR	16431	F3I	359-625
HEPATOCYTE GROWTH FACTOR PRECURSOR (SCATTER FACTOR) M60718	160718		
_		F3m	1549-19/0
HGE AGONIST/ANTAGOINST	146010	F3n	895-1051
COMPETITIVE HEPATOCYTE GROWTH FACTOR ANTAGONIST. AN M77227	177227		
ALTERNATIVE TRANSCRIPT OF THE HEPATOCYTE GROWTH			
FACTOR PRECURSOR (SCATTER FACTOR) (SF) (HEPATOPOEITIN A)			0007
		F4a	947-1908
IEN. GAMMA ANTAGONIST CYTOKINE	25270	F4b	395-685
	M27544; [M37484]		652-919
SI ELIKIN 1 RECEPTOR ANTAGONIST	M63099		225-1294
	M20566	F4e	2359-2823
	(02851	F4f	1107-1473
	(02770	F49	917-1208
	114844	F4h	181-436
SECTIBISOR (IL-3) (MULTIPOTENTIAL COLONY-	M14743; [M17115]		
CELL STIMULATING FACTOR) (MAST-CELL GROWTH FACTOR)			
(MCGE) (II.3)		F4i	390-608
4-I-4	M13982	F4j	216-459
-1	X04688; [J03478]	:	((
_		F4K	35-279
Œ	X04602; [M14584]		
		F41	130-555
	J04156	F4m	174-447
(P40)	X17543; [M30134]	F4n	156-399
	M57627	F5a	442-648
[adipogenesis inhibitory factor]	M57765	F5b	132-460
	M65291	F5c	066-009

TABLE 3 (CONT)

(NKSF,P40)	M65290 L06801	F5d	R22.848
	L06801		056-070
		F5e	285-743
	L15344	F5f	1181-1562
	U14407	F5g	338-695
INTERIFCIAL II-17	U32659	F5h	257-578
FERON ALPHA	J00209; [J00207]	F5i	89-430
	M28622	F5j	345-730
	X01992	F5k	391-586
E PEPTIDE	X02492	F5I	372-550
	X13967; [M63420]	F5m	1810-2239
M26	M25639	F5n	256-476
IBITE PROMOTING FACTOR(NEX!N), dia derived	A03911	F6a	667-915
OPHIC FACTOR)		492	112,416
WITH FACTOR 2) (NGF-2).	21.10	- -	21.7
NT-4 (NT-5) + NT-6 SA1	M86528; S41541; [541540; S41522	F6c	721-1079
	1141745	F6d	255-1326
PUGE ASSOC, PIONEIII	X06374	F6e	522-955
PRECURSOR	X02811; [X02744;		
	M12783]	F6f	1663-2125
	L36034	F6g	346-1241
	U16752; [L36033]	F6h	1053-1481
SELL FACTOR (C-KIT LIGAND)	M59964	F6i	898-1283
NO	M21626	F6j	273-504
	M96956; [M96955]		
(EPIDERMAL GROWTH FACTOR-LIKE CRIPTO PROTEIN CR1)			
(CRIPTO-1 GROWTH FACTOR) (CRGF) + TDGF2			
(TERATOCARCINOMA-DERIVED GROWTH FACTOR 2) (EPIDERIMAL		F6k	1294-1712
	L17075	F6I	814-1077
	J03241	F6m	
	L36052; [L36051; U11025]		
STIMULATING FACTOR) (C-MPL LIGAND) (ML) (MEGAKARYOCYTE		200	1416-1033
GDF) (THPO)			1410-1000
4	K03222	F/a	336-383
	X02812	F7b	2398-2575
CD27 (CD70 ANTIGEN)	L08096; [S69339]	F7c	233-627
CD30	L09753	F7d	627-1019
	L07414	F7e	863-1277

TABLE 3 (CONT)

Cell Cycle(Growth Begulators	GenBank #	Array Coordinate Position	Position
BETA CHAIN [Interferon gamma	U05875	F7!	1702-2039
receptor accessory factor-1 (AF-1) INTERFERON REGULATORY FACTOR [Interferon regulatory factor 1] X14454	X14454	F7g	478-695
¥	M91196	F7h	1253-1475
MINITED PROFESTION ALPHA-BETA RECEPTOR ALPHA-BETA RECEPTOR ALPHA-CHAINI	J03171	F7i	2562-2740
INTERFERON ALPHA-BETA RECEPTOR BETA CHAIN	X77722	F7	553-1012
INTERFERON GAMMA RECEPTOR ALPHA CHAIN	J03143	F7k	610-824
INTEREFRON GAMMA RECEPTOR	A09781	F7I	66-317
CAMMA INTERFERON INDITICED MONOKINE [Humia]	X72755	F7m	2021-2246
	X02530	F7n	280-613
HOUSEKEEPING GENES			

WO 98/53103 PCT/US98/10561

Apoptosis Array

5

In the apoptosis array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that are associated with apoptosis, e.g. cell cycle genes. In a specific apoptosis array of interest, the spots are as provided in Table 4.

TABLE 4

	ŀ	A area Considerate
GenBank #		Altay Cool umate
	CELL DIVISION CONTROL PROTEIN 2 HOMOLOG (EC 2.7.1) (P34 PHO LEIN	
X05360	KINASE) (CYCLIN-DEPENDENT KINASE 1) (CDK1)	38
	CELL DIVISION PROTEIN KINASE 2 (EC 2.7.1) (P33 PROTEIN KINASE)	Signal Control of the
	CELL DIVISION PROTEIN KINASE 3 (EC 2.7.1).	30
	1 .	3E
X66364	_	3F
		36
	CELL DIVISION PROTEIN KINASE 7 (EC 2.7.1) (CDK-ACTIVATING KINASE) (CAK)	
1 20320	(39 KD PROTEIN KINASE) (P39 MO15) (STK1) (CAK1).	ЭН
	CYCLIN-DEPENDENT KINASE 5 ACTIVATOR ISOFORM P391 PRECURSOR (CDK5	
1134051	ACTIVATOR) (P39I).	3
	CYCLIN-DEPENDENT KINASE 5 ACTIVATOR PRECURSOR (CDK5 ACTIVATOR)	
	(TAU PROTEIN KINASE II 23 KD SUBUNIT) (TPKII REGULATORY SUBUNIT) (P23)	
X80343	(P25) (P35).	37
	CDC25A; M.PHASE INDUCER PHOSPHATASE 1 (EC 3.1.3.48)	3K
18781871	CDC25B: M-PHASE INDUCER PHOSPHATASE 2 (EC 3.1.3.48). (CDC25HU2)	31.
	CDC25C: M-PHASE INDUCER PHOSPHATASE 3 (EC 3.1.3.48).	ЭМ
	CI K-1	3N
	CIK.2	30
	C X :3	48
	SEBINE/THREONINE-PROTEIN KINASE KKIALRE	4C
	SERINE/THREONINE-PROTEIN KINASE PCTAIRE-1	4D
	CEDINIC TUDE CONINE DE DE TEIN KINASE POTAIRE-2	4E
	SCONG THE CANADA PROTEIN KINASE POTAIRE.3	4F
1 25576	SERINE/THREONINE PROTEIN KINASE PITALRE	46
	CDC2-RELATED PROTEIN KINASE CHED	4H
1 33264	CDC2-RELATED KINASE PISSLRE	41
X51688	CYCLIN A	4.0
M26753	CYCLIN B1 G2/MITOTIC-SPECIFIC	4K
M74091	CYCLIN C G1/S-SPECIFIC	4L
X59798 [M64349]	CYCLIN D1 (CYCLIN PRAD1) (BCL-1 ONCOGENE)	4M
D13639 [M90813]	CYCLIN D2	4N
M92287	CYCLIN D3	40
M73812	CYCLINE	58
1147413 [] 49504]	CYC! IN G1	25

TABLE 4 (CONT)

		Array Coordinate
# 7 mg m	Colle Gene Name	Allay Cool Ciliane
Genbank #		50
U47414 [L49506]	N GZ	5E
J11791 [U12685]	CYCLIN H CYCLIN-DEPENDENT KINASE INHIBITOR 1 (MELANOMA DIFFERENTIATION ASSOCIATED PROTEIN 6) (MDA-6) (P21) (CDK-INTERACTING PROTEIN 1) (CIP1)	35
U09579; [L25610]	(WAF1) (CDKN1A) (CDKN1) (SDI1) (PIC1) (CAP2U) CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE	
J22398	INHIBITOR P57) (P57KIP2)	
197911	CYCLIN-DEPENDENT KINASE 4 INTELLIC COKNINA (MTS1). (CDKNINA) (MULTIPLE TUMOR SUPPRESSOR 1) (MTS1). (CDKNINA)	5H
- 1	CYCLIN-DEPENDENT KINASE 4 INHIBITOR B (P14-INK4B) (P15-INK4B) (MOL ITTLE	51
U17075; [L36844]	TUMOH SUPPHESSON 2) (MI 32) (SUS)	5.1
U40343; [U20498]	CYCLIN-DEFENDENT MINASE ASSEMBLY FACTOR MAT1 (RING FINGER PROTEIN	
	MAT1) (MENAGE A TROIS) (CDK7/CYCLIN H ASSEMBLY FACTOR) (P36) (P35	5K
X92669; [X87843]	(MNAT1) (MAT1) (CAP35).	51.
U10564	WEE1-LIKE PROTEIN KINASE (EC 2.7.1.12) (WEE1-LIK-1) (STPK13)	SM
U01038	SERINE/THREONINE-PHOLE IN NIMBER TEN LEG ENTRY AND AND AND AND AND AND AND AND AND AND	NS.
U38545	PHOSPHOLIPASE U1	50
D63878	NEDDS PROTEIN HOMOLOG.	89
872008	CDC10 PROTEIN HOMOLOG	29
	CDC27HS PHOLEIN	Q9
122005	UBIQUITIN-CONJUGATING ENGINE EX-COOST	6E
	CDC16HS.	6F
	CDC37 HOMOLOG.	98
U77949	CDC6-RELATED PHOTEIN	
	EXTHACELLULAH SIGNAE-TICOCCITE (MAPK 1) (P44-ERK1) (ERT2) (P44-ERK1) (STIMULATED MAP2 KINASE) (MAPK 1) (MAPK 1) (MAPK 1) (FA4-ERK1) (ERT2) (P44-ERK1)	Н9
X60188	MAPK) (MICROTUBULE-ASSOCIALED PHOLEIN-S MINASE) (MITOGEN	
	EXTRACELLULAR SIGNAL-HEGULA I EU MINASE 2 (EO 2:11:1) (ETT1).	19
M84489	EXTRACELLULAR SIGNAL-REGULATED KINASE 3 (EC 2.7.1) (ERK3) (MAP	9
X80692	KINASE ISOFORM P97) (P97-MAPK).	
	EXTRACELLULAR SIGNAL-HEGULATED MINASE + (EX 2.7.1.1.) (E. 11.7.) (MIN. 11.)	9 QK
X59727	EXTRACELLULAR SIGNAL-REGULATED KINASE 5 (EC 2.7.1) (ERKS) (ERK4)	- P
U25278	(BMK1 KINASE)	

TABLE 4 (CONT)

		Array Coordinate
GenBank #		W9
X79483		
	MITOGEN-ACTIVATED PROTEIN KINASE 1738 (EC 2.7.1.1) (MAT NINASE 1 33) (CYTOKINE SUPPRESSIVE ANTI-INFLAMMATORY DRUG BINDING PROTEIN)	
		N9
L35253; [L35263]	KINASE MXIZ).	
	KINDAR 1) (INK-46)	09
20310	STRESS-ACTIVATED PROTEIN KINASE JNK2 (EC 2.7.1) (C-JUN N-TERMINAL	7B
L31951	KINASE 2) (JNK-55).	
	STRESS-ACTIVATED PROTEIN KINASE JUKS (EC Z.C.1.7) (C-3014 IV-1.5) INVINCED PROTEIN KINASE DAG 2619)	7C
U34819; [U07620]	KINASE 3) (JUKS) (MAT NINASE 143 ST. 12): D) IA SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 5 (EC 2.7.1)	
1125265	(MAP KINASE KINASE 5) (MAPKK 5) (MAPK/ERK KINASE 5).	a)
05050	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE I (EC. 2.7.1.7)	
	(MAP KINASE KINASE 1) (MAPKK 1) (EHK ACTIVATOR NINASE 1) (MAPKK 1)	ŻΕ
L05624	KINASE 1) (MEK1).	
	DUAL SPECIFICITY MITOGEN-ACTIVALED PROTEIN MINASE 6: (SAPKK3)	
U39657	(MAP KINASE KINASE O) (MAP KINASE KINASE O)	76
U78876	MEK KINASE 3	H/
M15796; [J04718]	PCNA (CYCLIN)	12
U49070	PIN1	
	A-ASSOCIATED PROTEIN	L2
M15400	SUSCEPTIBILITY	7K
X74594	RB2/P130	7.
X74262	RBA/P48	J.W.
S66431	HBP2 HELINOBLASTOMA-BINDING PROTEIN)	N/
55/153; 55/160	DECA DETINION ASTOMA BINDING PROTEIN	70
X85133		8B
X85134	HBQ-3	8C
M96577	E2F-1 PHB-BINDING Thousand	08
Y10479	E2F-3	8E
U15642	E2F-5	8F
123959	E2F-RELATED TRANSCHIP ION FACTOR (Dr-1)	86
U18422	DP2 (HUMDP2), DIMERIZATION PARTINER OF EST	BH.
U23435; U31089	ABL INTERACTOR 2 (ABI-2) + ABL BINDING FROTEIN 3 (ABLE) 3) (1111)	81
129511	GRB2 (GROWIH FACTOR RECEPTION-BOOKS TROUBLES E	

TABLE 4 (CONT)

GenBank #	Cell Cycle - Gene Name	Array Coordinate	
U69276	GRB-IR / GRB10	8.1	
X03484	RAF ONCOGENE	8K	
M95712	RAF,B-	8L	
	TRANSCRIPTION FACTOR AP-1 (C-JUN PROTO ONCOGENE)	8M	
	JUN B TRANSACTIVATOR	8N	
	TRANSCRIPTION FACTOR JUN-D	80	
	N-MYC	98	
D89667		26	
	NUCLEOSIDE DIPHOSPHATE KINASE B (C-MYC TRANSCRIPTION FACTOR		_
	(PUF)	ЭD	
X16416 [M14752]	c-abi	3 6	
	р53 РАТНWAY		
M14694	CELLULAR TUMOR ANTIGEN P53	9F	
	MDM2 PROTEIN (PS3-ASSOCIATED PROTEIN) + MDM2-A (GB: U33199) + MDM2-C		
Z12020	(GB: U33201)	56	
AF007111	MDM2-LIKE P53-BINDING PROTEIN (MDMX)	Н6	
Y11416	P73, A MONOALLELICALLY EXPRESSED P53-RELATED PROTEIN	16	
AF010310 AF010311	AF010310 AF010311 P53 INDUCED PROTEIN	P)	
	PiG3 (PiG3)	9K	
	PIG7 (PIG7)	J6	
AF010314	PIG10 (PIG10)	9M	
AF010315	PIG11 (PIG11)	N6	
AF010316	PIG12 (PIG12)	06	
U90313	GLUTATHIONE-S-TRANSFERASE HOMOLOG	10B	
U66469	P53-DEPENDENT CELL GROWTH REGULATOR CGR19	10C	
AF001954	GROWTH INHIBITOR P33ING1 (ING1)	10D	
L13698	GROWTH-ARREST-SPECIFIC PROTEIN 1 (GAS-1).	10E	
	BCL FAMILY		
	BCL2	10F	
	BCL2 AND P53 BINDING PROTEIN BBP/53BP2 (BBP/53BP2)	10G	
	BAX	10H	
U59747	APOPTOSIS REGULATOR BCL-W	101	
L08246	INDUCED MYELOID LEUKEMIA CELL DIFFERENTIATION PROTEIN MCL-1 (ORF IS AT NT. 61-1053; ML)	107	

TABLE 4 (CONT)

		Array Coordinate	
GenBank #	RLY	, to	
		10L	
V80086- [1]34584]		10M	
1123765 FU16812; U	BCL-2 HOMOLOGOUS ANTAGONIST/KILLEH (APUT 10313 11ECCE	10N	
1	BRAG-1=BRAIN-RELATED APOPTOSIS GENE/BUL-2 NUMOLOG	100	
1 1	BAD PROTEIN (BCL-2 BINDING COMPONENT B). BCL-2 BINDING ATHANOGENE-1 (BAG-1) (GLUCOCORTICOID RECEPTOR-	118	
S83171: [Z35491]	ASSOCIATED PROTEIN RAP46).	11C	
U76376	Harakiri, a protein that activates cell deaut and inclass.		
	CASPASE CASCADE		
	CASPASES CASPASES	110	
1113699 IM87507; X6(ICE)	G(ICE) (INTERLEUKIN-1 BETA CONVEHI ING ENCTIME) IL 13/10:00	11E	
U13021; [U13022]	(CASPASE-2) (ICH-1L) (ICH-1S) APOPAIN PRECURSOR (EC 3.4.22) (CYSTEINE PROTEASE CPP32) (YAMA APOPAIN PRECURSOR (EC 3.4.22) (CYSTEINE PROTEIN) (CASPASE-3) ISOFORM	u.	
1113737	ALPHA		
	ICH-2 PROTEASE PRECURSOR (EC 3.4.22) (1.4 PROTEASE) (CASPASE-4) + CASPASE-5 PRECURSOR (EC 3.4.22) (ICH-3 PROTEASE) (TY	116	
U28014; U28015	PROTEASE ((CEREL-III).	-	
	CASPASE-6 PHECONOCIA (EO CITETA) (CASPASE-6 PHECONOCIA (EO CITETA) (CASPASE-6 PHECONOCIA (CASPASE-6 PHECONOC	11H	
U20537; U20536	ISOFORM BETA + ISOFORM ALT TO (ICE-LIKE APOPTOTIC PROTEASE 3) (ICE-CASPASE-7 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC PROTEASE 3) (ICE-	11	
U37448	LAP3) (APOPTOTIC PROTEASE MCH-3) (CMH-1) (LIOES)		
	CASPASE-8 PRECURSON (EC 3:4.62.7) (136.6.4.1) (FADD-HOMOLOGOUS IMPORT1-ASSOCIATED CED-3 HOMOLOG) (MACH.) (FADD-HOMOLOGOUS		
	ICE/CED-3-LIKE PROTEASE) (FADD-LIKE ICE) (FLICE) (APOFTOTICE OF STEINE) ICE/CED-3-LIKE PROTEASE MCH.S.) (CAPA) (CASP8) (MCH5) ISOF	113	
U60520; U58143; X98PRC	GASPASE-8 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC PROTEASE 5)		
	(MORTI-ASSOCIATED CED-3 HOMOLOG) (MACH) (FADD-100000000000000000000000000000000000	7	
1150520: 1158143: >	HENESON 1158143" X98 PROTEASE) (APOPTOTIC PROTEASE MCH-5) (CAP4) (CASP8) (MCH5) ISON	+-	
0000000	CASPASE-9 PRECURSOR (EC 3.4.22) (ICE-LIKE APOPTOTIC PROTECTED 5)	111	
U56390; [U60521]	LAP6) (APOPTOTIC PHOTEASE MOTE) (EC 3.4.22) (APOPTOTIC ICE-LIKE APOPTOTIC PROTEASE 4 PRECURSOR (EC 3.4.22) (APOPTOTIC	11M	
U60519	PROTEASE MCH-4) (CASPASE-10)		

TABLE 4 (CONT)

GenBank #	Cell Cycle - Gene Maine	
141690	D PROTEIN (TRADD)	110
	_	12B
[L81153]	TRAF6	
		12C
U59863; [U63830]	THAE INTERACTING PROTEIN (TRIP)	12D
V10256	SERINE/THREONINE PROTEIN KINASE, NIK; BINDS SPECIFICALLY TO TRAF2	12E
00001	CASPER, A FADD. AND CASPASE-RELATED INDUCER OF APOPTOSIS ICASH-	12F
AF010127[Y14039; Y	ALPHA+ CASH-BE IAI (FLAME: 1) IL LIOU LINE IN THE STATE ADAPTOR	
	MOLECULE FOR CASPASE-2 AND FASI/TNF RECEPTOR-INTERACTING PROTEIN	12G
U84388	RIP	
U25994; [U50062]	CELL DEATH PROTEIN KINASE HILL CELL DEATH PROTEIN THAT ACTIVATES INK AND APOPTOSIS	121
AF015956	DAXX, A FAS-BINDING FROTEIN TITLE OF SOCIATED PROTEIN (TRAP3)	
1		123
012597	U1259/ CD40 RECEPTOR ASSOCIATED FACTOR 1 (CRAF1) (CAP-1), (LMP1 ASSOCIATED	10%
1121092- [U15637; L3	PROTEIN)	151
	INHIBITOR OF APOPTOSIS PROTEIN 1 (HIAP1) (HIAP-1) (C-IAP2) (1NFR2-1 PAF	12L
U45878; [U37546]	SIGNALLING COMPLEX PHOLEIN 1) (IAT HOMOCOCO) (10.18P1) (TNFR2- TRAF	
	INHIBITOR OF APOLITIONS FIRST ENTIRE J. (MINE).	12M
U45879; IU37547J	X-LINKED INHIBITOR OF APOPTOSIS PROTEIN (X-LINKED IAP) (IAP-	Not
U45880; [U32974]	LIKEPROTEIN) (HILP).	
	LICANDS AND BECEPTORS	
70400	[TNF-a]	120
701394	LYMPHOTOXIN-ALPHA FORMERLY TUMOR NECROSIS FACTOR BETA (TNF-	- 14B
D12614	b)]	14C
L11015	LYMPHOTOXIN-BETA	14D
U69611	TNF-ALPHA CONVEHING ENZINGE EAS ANTIGEN I IGAND (APOPTOSIS ANTIGEN LIGAND) (APTL) (APT1LG1) (FASL)	
D20429- [1108137]		14E
U38122, [U00137]	THE STATES APOBIOSIS INDIVING LIGAND TRAIL)	14F

TABLE 4 (CONT)

	- 1	Array Coordinate
#	Cell Cycle - Gene Name	146
	1 / 1	14G
	SECRETED APOPTOSIS RELATED PROTEIN 3 (SARP3)	14H
	TUMOR NECROSIS FACTOR RECEPTOR TUMOR NECROSIS FACTOR	158
M33294	TUMOR NECROSIS FACTOR RECEPTOR (TUMOR NECROSIS FACTOR	C
M32315	RECEPTOR 2]	130
	FAS/APO 1	156
	CYTOTOXIC LIGAND I MAIL MECENTON	15F
AF016268 DEA	DEATH RECEPTOR 5 (DHS)	15G
109392, 1073300,074	II IN I IKE GROWTH FA	15H
	INCOLINATIVE GROWTH FACTOR II Somatomedin A)	16B
WC3043	INSUITING GROWTH FACTOR I RECEPTOR	16C
	CATION-INDEPENDENT MANNOSE-6-PHOSPHATE RECEPTOR [insuline-like	160
[103528]	growth factor receptor ii, furth-z	16F
	IGFBP COMPLEX ACID LABILE CHAIN	120
M35410	IGFBP2	
	IGEBP3 (GROWTH HORMONE-DEPENDENT INSULIN-LINE GROWIN FACTOR:	16G
M358/8]	BINDING FROIEIN	16H
M62403	Lorent 4	178
Mosnoz		17C
M62402	IGHBP6	
	OTHER REGIII ATORS	
	DEATH-ASSOCIATED PROTEIN 3 (DAP-3) (ionizing radiation resistance conferring	
U18321; [X83544]	protein)	מינ
X76104	DEATH-ASSOCIATED PROTEIN KINASE 1 (EC 2.7.1) (DAP KINASE 1).	1/E
X86779	Fas-activated serine/threonine kinase (FAST) phosphorylates TIA-1	1/1
S78085	PDCD2	1/6
M63167	Akt1 (rac protein kinase alpha, protein kinase B, c-Akt)	17H
M77198; [M95936]	AKT2 (rac protein kinase beta)	188
U63295	seven in absentia homolog	18C
1137688	PATS1	18D
1191985	DNA fragmentation factor-45	18E
AF022385	apoptosis-related protein TFAR15 (TFAR15)	18F
0.000	The description of the state of	18G

TABLE 4 (CONT)

GenBank #	Cell Cycle - Gene Name	Array Coordinate
182938		18H
1122286	chromosome segregation gene homolog CAS	19B
033200	anotocis inhibitor survivin	19C
0/200	OTD hisding protein (tho A)	19D
722080	GIT-Diffailing protein (filed)	19E
L09210	MILHIC OXIDE STRIFFASE (ZA, INDUCIBLE)	196
M58603		100
M83221	TRANSCRIPTION FACTOR RELB [I-Hel]	0.00
U08015	NF-ATc [Transcription factor (NFATc.b)]	Z05
D15057	DAD-1 [DEFENDER AGAINST CELL DEATH 1]	SOC
	CLUSTERIN [complement lysis inhibitor; testosterone-repressed prostate message 2;	
M74816	apolipoprotein J; sulfated glycoprotein-2	2002
D13889	DNA-BINDING PROTEIN INHIBITOR ID-1	20E
X15722	GLUTATHIONE REDUCTASE	ZUF
J03746	TATHIONE S-TRANSFERASE MICRO	20G
	GLUTATHIONE S-TRANSFERASE M4 [GLUTATHIONE S-TRANSFERASE MU 1]	
X08020		218
X15480	TATHIONE S-TRANSFERASE P	212
	GLUTATHIONE S-TRANSFERASE A1-1 (Glutathione S-transferase (GST) Ha	
M14777	subunit 1)	710
M21304	GLUTATHIONE PEROXIDASE	712
X79389	GLUTHATHIONE S-TRANSFERASE (THETA 1)	21F
590469	NADPH-CYTOCHROME P450 REDUCTASE	21G
	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD153 (DNA-	800
S40706 [S62138]	DAMAGE INDUCIBLE PROTEIN) (CHOP).	22B
	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PHOLEIN GADD45 (DNA-	200
M60974	DAMAGE INDUCIBLE I HANSCHIPT 1) (DOITT).	220
U15172	NIPI	29E
U15174	NIP3	22F
L07414	1	220
960807	CD27 LIGAND (CD70 antigen)	000
X96586	FAN PROTEIN	230
M84820	RETINOIC ACID RECEPTOR RXR-BETA	23C
X07282	RETINOIC ACID RECEPTOR BETA-2	23D
M93426	PROTEIN-TYROSINE PHOSPHATASE ZETA	23E
	PROPERTY OF THE PROPERTY EDUCE	235

TABLE 4 (CONT)

		Array Coordinate
GenBank #	Cell Cycle - Gene Name UV EXCISION REPAIR PROTEIN PROTEIN RAD23 [xeroderma pigmentosum group UV EXCISION REPAIR PROTEIN PROTEIN RAD23 [xeroderma pigmentosum group	23G
D21090	C repair complementing protein poorningobl	
-	SOURCE TO THE SENE CONTRACTOR OF THE SENE CON	
1	HOUSEKEEPING GENES	1A
	UBIQUITIN	18
	PHOSPHOLIPASE AZ	10
	HYPOXANTHINE-GUANINE PROSTHONDOS ENTRACE	10
	GLYCERALDEHYDE 3-PHOSPITATE DETTOGETHER	16
	TUBULIN ALPHA	4
	HLA CLASS I HISTOCOMPATIBILITY ANTICEN, OFFICE TO THE CLASS I HISTOCOMPATIBILITY ANTICENS.	16
	BETA-ACTIN	Ŧ
	23 kD HIGHLY BASIC PROTEIN	
	RIBOSOMAL PROTEIN S9	
	NEGATIVE CONTROLS	1.3
	M13 mp18(+) STRAND DNA	1 x
	I-DNA	
	pUC 18	
		1M1N1O1P
	CALIBRATION MAHKEHS	
	ORIENTATION MAHKEHS	2D2G2J2M3A3P6A6P9A9P12A12
	Dark spots	2A2B2C2E2F2H2I2K2L2N2O2P4/
	Faint spots	
	Column 13 is plank	

WO 98/53103 PCT/US98/10561

Human Stress Array

In the human stress array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that are associated with stress responses of human cells, e.g. stress response regulators and effectors. In a specific human stress array of interest, the spots are as provided in Table 5.

TABLE 5

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
K00650	C-fos
	CAMP RESPONSE ELEMENT BINDING PROTEIN CRE-BP1 (CAMP responsive element binding protein 1)
M34356	CREB (ACTIVE TRANSCRIPTION FACTOR)
X60188	EXTRACELLULAR SIGNAL-REGULATED KINASE 1 (EC 2.7.1) (ERK1) (INSULIN- SIIMULAIED MAP2 KINASE) (MAP KINASE 1) (MAPK 1) (P44-ERK1) (ERT2) (P44-MAPK) (MICROTUBULE-ASSOCIATED PROTEIN-2 KINASE).
M84489	EXTRACELLULAR SIGNAL-REGULATED KINASE 2 (EC 2.7.1) (ERK2) (MITOGEN- ACTIVATED PROTEIN KINASE 2) (MAP KINASE 2) (MAPK 2) (P42-MAPK) (ERT1).
X80692	EXTRACELLULAR SIGNAL-REGULATED KINASE 3 (EC 2.7.1) (ERK3) (MAP KINASE ISOFORM P97) (P97-MAPK).
X59727; S38873	EXTRACELLULAR SIGNAL-REGULATED KINASE 4 (EC 2.7.1) (ERK4) (MAP KINASE ISOFORM P63) (P63-MAPK).
U25278	EXTRACELLULAR SIGNAL-REGULATED KINASE 5 (EC 2.7.1) (EKKS) (EKK4) (BINK) I NINASE).
X79483 U53442	EXTRACELLULAR SIGNAL-REGULATED KINASE 6 (EC 2.7.1) (ERK6) (ERK5). MITOGEN-ACTIVATED PROTEIN KINASE P38 BETA (EC 2.7.1) (MAP KINASE P38 BETA).
126318	STRESS-ACTIVATED PROTEIN KINASE JNK1 (EC 2.7.1) (C-JUN N-TERMINAL KINASE 1) (JNK
131951	46) STRESS-ACTIVATED PROTEIN KINASE JNK2 (EC 2.7.1) (C-JUN N-TERMINAL KINASE 2) (JNK- 55).
U25265; (U71087; U71088)	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 5 (EC 2.7.1)(MAP KINASE KINASE 5) (MAPKK 5) (MAPK/ERK KINASE 5) (MEK5)
	MAP KINASE KINASE MEK5B. MAP KINASE KINASE MEK5C

TABLE 5 (CONT)

,	
GenBank#	STRESS RESPONSE REGULATORS AND EFFECTION STRESS RESPONSE REGULATORS AND EFFECTION (MAP
105624	DUAL SPECIFICITY MITOGEN-ACTIVATOR KINASE 1) (MAPK/ERK KINASE) (MEK1). KINASE KINASE 1) (MAPKK 1) (ERK ACTIVATOR KINASE 1) (MAPK/ERK KINASE)
111285	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 2 (EC 2.7.1.) (MAP) KINASE 2) (MAPK/ERK KINASE) (MEK2). KINASE KINASE 2) (MAPK/ERK KINASE) (MEK2).
U39657	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE 6 (EC 2.7.1)(MAPKINASE 6) (SAPKK3).
U78876	MEK KINASE 3 STESSLIKE KINASE OXIDANT STRESS KINASE (YSK1, STEZO and SPS1 RELATED KINASE)
D6378U	SPS1/STE20 HOMOLOGUE, KHS, ACTIVATOR OFJUN N-TERMINAL KINASE (HSU77129)
1107340	B LYMPHOCYTE GERMINAL CENTER KINASE (HSU07349)
U66464	HEMATOPOIETIC PROGENITOR KINASE ACTIVATOR OF STATES ACTIVATOR OF STATES ACTIVATION OF STATES
AB005216	NCK, ASH AND PHOSHPHOUPASE C GAMMA-BINDING PROTIEN NAP4(AB005216) NCK MELANOMA CYTOPLASMIC SRC HOMOLOGUE (HSNCK)
U24153	SERINE/THREONINE-PROTEIN KINASE PAK-GAMMA (EC 2.7.1) (GAMMA-PAK) (P21-ACTIVATED KINASE 3) (PAK65) (S6/H4 KINASE) (PAK2) PAK3.
M35543	G25K GTP-BINDING PROTEIN, BRAIN ISOFORM (GP) (CDC42 HOMOLOG) CDC42.
U12595	TUMOR NECROSIS FACTOR TYPE 1 RECEPTOR ASSOCIATED TO SECURITION OF SECURI
012596	TUMOR NECROSIS FACTOR TYPE 1 RECEPTOR ASSOCIATED PROTEIN(1) (1) TO THE TOTAL AS ALICE KINASE A) (1) MOR
X17620	NUCLEOSIDE DIPHOSPHATE KINASE A (EC 2.7.4.6) (NDK A) (NDP NIMASE Y) (NOT NIMASE A (EC 2.7.4.6) (NDE ASTATIC PROCESS-ASSOCIATED PROTEIN) (METASTASIS INHIBITION FACTOR NIM23)
	(NM23-H1). (NM23-H1). (HSF 1) (HEAI SHOCK TRANSCRIPTION FACTOR 1)(HSTF
M64673	HEAL SHOCK FACTOR INCIDENT A CHEAT SHOCK TRANSCRIPTION FACTOR 2)(HSIF
M65217	HEAT SHOCK FACTOR PROTEIN 2 VISI 2) VISI 1.
D87673	HEAT SHOCK TRANSCRIPTION FACTOR 4. FKBP-RAPAMYCIN ASSOCIATED PROTEIN (FRAP) (HUMFRAPX)
1000	

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
M35663: (U50648)	INTERFERON-INDUCIBLE RNA-DEPENDENT PROTEIN KINASE (P68 KINASE)
	10 KD HEAT SHOCK PROTEIN, MITOCHONDRIAL (HSP10) (10 KD CHAPERONIN)
D86956	HEAT-SHOCK PROTEIN 110 KD (KIAAD201)
X54079; (x03900,L39370;	HEAT SHOCK 27 KD PROTEIN (HSP 27)(STRESS-RESPONSIVE PROTEIN 27)(SRP27)(ESTROGEN-REGULATED 24 KD PROTEIN) (28 KD HEAT SHOCK PROTEIN).
X16477; Z23090; S74571)	
D83174	47 KD HEAT SHOCK PROTEIN PRECURSOR (COLLAGEN-BINDING PROTEIN 1) (COLLIGIN 1)
	Collagen binding protein 2 (HUMCBP2).
M11717; (M59828)	M11717; (M59828) HEAT SHOCK 70 KD PROTEIN 1 (HSP70.1) (HSP70-1/HSP70-2).
126336	HEAT SHOCK-RELATED 70 KD PROTEIN 2 (HEAT SHOCK 70 KD PROTEIN 2).
L12723	HEAT SHOCK 70 KD PROTEIN 4 (HSP/AKY). HEAT SHOCK 70 KD PROTEIN 6 (HEAT SHOCK 70 KD PROTEIN B').
A31/37, M11230	HEAT SHOCK 70 KD PROTEIN 7 (HEAT SHOCK 70 KD PROTEIN B) (FRAGMENT).
Y00371	HEAT SHOCK COGNATE 71 KD PROTEIN.
X07270; (X15183;	HEAT SHOCK PROTEIN HSP 90-ALPHA (HSP 86).
M27024; M30626;	
M3062/)	CO GOAL VISION OF BETA A HED BOT HISTORY WOOL STATE
M16660 U15590	HEAT SHOCK PROTEIN TO? YEACH) HEAT SHOCK PROTEIN 27 (heart)
267070	HEAT SHOCK PROTEIN HSP72 HOMOLOG (FRAGMENT).
U40992	HEAT SHOCK PROTEIN HSP40HEAT SHOCK PROTEIN HSP40 HOMOLOG.
115189	REGULATED PROTEIN) (GRP 75) (PEPTIDE-BINDING PROTEIN 74) (PBF74) (MOKHLIN) (MOD).
U28918	HSC70-INTERACTING PROTEIN (PROGESTERONE RECEPTOR-ASSOCIATED P48 PROTEIN)
D13388	DNAJ PROTEIN HOMOLOG 2 (DNAJ2 OR HDJ2)
D49547; (D17749; D85429)	
M19645	78 KD GLUCOSE REGULATED PROTEIN PRECURSOR (GRP 78) (IMMUNOGLOBULIN HEAVY CHAIN BINDING PROTEIN) (BIP)

TABLE 5 (CONT)

	CONTRACTOR OF THE PERSONS
GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
L10284; (L18887; M94859; M98452)	CALNEXIN PRECURSOR (MAJOR HISTOCOMPATIBILITY COMPLEX CLASS I ANTIGEN- BINDING PROTEIN P88) (P90) (IP90)
1	CALRETICULIN PRECURSOR (CRP55) (CALREGULIN) (HACBP) (ERP60)(52 KD RIBONUCLEOPROTEIN AUTOANTIGEN RO/SS-A)
305016	PROTEIN DISULFIDE ISOMERASE-RELATED PROTEIN PRECURSOR (ERP72)
(1.24805)	P23 PROGESTERONE RECEPTOR ASSOCIATED PROTEIN (HUMPRA)
M86752	TRANSFORMATION -SENSITIVE PROTEIN (IEF SSP 3521)
111667	CYCLOPHILIN-40
U73704	48 kDa FKBP-ASSOCIATED PROTEIN FAP48
U42031	54 KDA PROGESTERONE RECEPTOR-ASSOCIATED PROTEIN FKBP54
M34539; (M80199; M80706;M92423; IO5340; X55741;	M34539; (M80199; FK506-BINDING PROTEIN (FK8P) (FK8P12) (PEPTIDYL-PROLYL CIS-TRANS ISOMETASE) M80706;M92423; (PPIASE) (ROTAMASE) IN5340: X55741:
x52220)	
M88279	IMMUNOPHILLIN (FKBP52)
M65128	RAPAMYCIN-BINDING PROTEIN (FKBP-13)
X56134 (M14144;	VIMENTIN, INTERMEDIATE FILAMENT PROTEIN
M34664; (M22382	M34664: (M22382) MITOCHONDRIAL MATRIX PROTEIN PI PRECURSOR (P60 LYMPHOCYTE PROTEIN) (HSPD1 OR HSP60) (CHAPERONIN HOMOLOG) (HUCHA60) (HEAT SHOCK PROTEIN 60)
583171; (235491)	BCL-2 BINDING ATHANOGENE-1 (BAG-1) (GLUCOCORTICOID RECEPTOR-ASSOCIATED PROTEIN PAPAS)
D23662	UBIQUITIN-LIKE PROTEIN (NEDD8)
X52882	T-COMPLEX PROTEIN 1, ALPHA SUBUNIT (TCP-1-ALPHA)(CCT-ALPHA) CCT1 OR CCTA OR TCP1
U38846	1-COMPLEX PROTEIN 1, DELTA SUBUNIT (TCP-1-DELTA)(CCT-DELTA) (STIMULATOR OF TAR RNA BINDING) (HSU38846).
D43950	1-COMPLEX PROTEIN 1, EPSILON SUBUNIT (TCP-1-EPSILON)(CCT-EPSILON) (HUMKG1DD)
X74801; (U17104)	1-COMPLEX PROTEIN 1, GAMIMA SUBUNIT (TCP-1-GAMIMA)(CCT-GAMIMA) (CCT3) OR (CCTG) OR (TRIC5) (HSHUMAPC).

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
U83843	T-COMPLEX PROTEIN 1, ETA SUBUNIT (TCP-1-ETA) (CCT-ETA)(HIV-1 NEF INTERACTING PROTEIN) (HSUB3843).
013627	T-COMPLEX PROTEIN 1, THETA SUBUNIT (TCP-1-THETA)(CCT-THETA) (HUMRSC548).
	HEME OXYGENASE 1 (EC 1.14.99.3) (HO-1) (HSOXYGR).
(\$34389)	HEME OXYGENASE 2 (EC 1.14.99.3) (HO-2)
۱_	ENDOPLASMIN PRECURSOR (94 KD GLUCOSE-REGULATED PROTEIN) (SK779) (SK779) (HOMOLOG) (TUMOR REJECTION ANTIGEN 1) (HSTRA1).
005569	ALPHA CRYSTALLIN A CHAIN (HSU06569).
\$45630	ALPHA CRYSTALLIN B CHAIN (ALPHA(B)-CRYSTALLIN) (ROSENIHAL FIBER COMPONENT).
U59058	BETA CRYSTALLIN A3 (HSU59058).
U59057	BETA CRYSTALLIN A4 (HSU59057).
U35340	BETA CRYSTALLIN B1 (CRYBB1) (HSU35340).
110035	BETA CRYSTALLIN B2 (BP) (HUMCRYB2B).
071216	BETA CRYSTALLIN BS (VCKTBBS OR CATES) (1805) (E19)
136869	BETA CRYSTALLIN S (GAMINIA CRYSTALLIN S) (CRYTOS) CR (CRYTOS)
U66582; M11971;	GAMMA CRYSTALLIN C (GAMMA CRYSTALLIN 2 OR 1/3) (CRYGC) OR (CRYG3).
(M11970)	CENTER OF CONTRACTOR 1-3 (CRYGR) OR (CRYGS)
	(HUMCRYGX1).
102950	MU-CRYSTALLIN HOMOLOG (CRYM) (HUMMUCRYS).
L13278; (S58039)	QUINONE OXIDOREDUCTASE (EC 1.6.5.5) (NADPH: QUINONE REDUCTASE) (ALT.)
D16234 749835	PROBABLE PROTEIN DISULFIDE ISOMERASE ER-60 PRECURSOR (EC 5.3.4.1) (ERP60)
D83485; U42068)	(58KDA MICROSOMAL PROTEIN) (phospholipase C-alpha)
D49489	PROTEIN DISULFIDE ISOMERASE PS PRECUISOR (EC 5.3.4.1) (HUMP3).
M75715	EUKARYOTIC PEPTIDE CHAIN RELEASE FACTOR SUBUNIT 1 (ERF.1) (185-1) (CT.1-1-NOTE.19) [RF1.
D49490	PROTEIN DISULFIDE ISOMERASE-RELATED PROTEIN PRECURSOR (EC 5.3.4.1) (PDIR) (HUMPDIR).
J02783: (X05130;X07077)	
	(P55)(HSPRO4HY).

TABLE 5 (CONT)

(Glu (AB6737 STE X63368; (S37374; DN S37375) 15	Chitathionaling transhydrogenase (EC 5.3.4.) / 1.8.4.2); protein-disulfide reductase
(537374;	(glutathione) (HSGITR).
(537374;	STRUCTURE-SPECIFIC RECOGNITION PROTEIN 1 (SSRP1) (RECOMBINATION SIGNAL SEQUENCE RECOGNITION PROTEIN) (T160) SSRP1.
	DNAJ PROTEIN HOMOLOGS HSJ1A protein; HSJ1B protein.(HSJ-1)(HSHSJ1MIt)
	150 KDA OXYGEN-REGULATED PROTEIN ORP150 (HSU65785)
NO	DNA DAMAGE RESPONSE/REPAIR/RECOMBINATION
: (L40817;	MUSCLE-SPECIFIC DNASE I-LIKE (DNase X) (XIB)
124544 (RAD	
4	TRANSCRIPTIONAL ACTIVATOR PROTEIN PUR-ALPHA
M29971 ME	METHYLATED-DNAPROTEIN-CYSTEINE METHYLTRANSFERASE (6-O-METHYLGUANINE-DNA METHYLTRANSFERASE) (MGMT)
U09579; (1.25610) CY PR (C)	CYCLIN-DEPENDENT KINASE INHIBITOR 1 (MELANOMA DIFFERENTIALION ASSOCIALED PROTEIN 6) (MDA-6) (P21) (CDK-INTERACTING PROTEIN 1) (CIP1) (WAF1) (CDKN1A) (CDKN1) (SD1) (PIC1) (CAP20)
L37374 FL	FLAP ENDONUCLEASE-1 (MATURATION FACTOR 1) (MF1) (FEN-1)
U70310 DI	DNA REPAIR PROTEIN XRCC9
HT3218 (X02317; SU	SUPEROXIDE DISMUTASE (CU-ZN) (EC 1.15.1.1) SOD1.
	EXTRACELLULAR SUPEROXIDE DISMUTASE PRECURSOR (CU-ZN) (EC 1.15.1.1) (EC-SOD) SOD3
X07834; (X59445) St	SUPEROXIDE DISMUTASE PRECURSOR (MIN) (EC 1.15.1.1) SOD2
M14694; (M14695) C	M14694; [M14695] CELLULAR TUMOR ANTIGEN P53
Z12020; (M92424) N	212020; (M92424) MDM2 PROTEIN (P53-ASSOCIATED PROTEIN)
2 2	MDM2-C (GB: U33201)
	ATAXIA TELANGIECTASIA (ATM)
J03250 D	DNA TOPOISOMERASE I (TOP1) DNA TOPOISOMERASE II, ALPHA (TOP2A)
	DNA TOPOISOMERASE II, BETA (TOP2B)
	DNA TOPOISOMERASE III (TOP3)

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
\$40706 (\$62138)	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD 153 (DNA-DAMAGE INDUCIBLE TRANSCRIPT 3) (DDIT3) (C/EBP-HOMOLOGOUS PROTEIN) (CHOP)
X04076	CATALASE (EC 1.11.1.6) CAT.
X51420	5.6-DIHYDROXYINDOLE-2-CARBOXYLC ACID OXIDASE PRECURSOR (DHICA OXIDASE) (TYROSINASE-RELATED PROTEIN 1) (TRP-1) (CATALASE B) (GLYCOPROTEIN-75) (GP75)
	BASE EXCISION REPAIR
X15653	URACIL-DNA GLYCOSYLASE PRECURSOR (UNG.1)
X52486	URACIL-DNA GLYCOSYLASE 2 (UNG2)
M74905	DNA-3 METHYLADENINE GLYCOSYLASE (3-METHYLADENINE DNA GLYCOSYLASE)
	(ADPG) (3-ALKYLADENINE DNA GLYCOSYLASE) (N-WEITH DANNE CAN COLONG CONTROL (MAG I) (3MeAG)
051166	G/T MISMATCH-SPECIFIC THYMINE DNA GLYCOSYLASE (TDG)
Y11838	8-OXYGUANINE DNA GLYCOSYLASE HOMOLOG I (mutim HOMOLOG) (UGHI)
	(HOGGI) (Farye)
U63329	muty HOMOLOG (HMYH)
	SAA JOHN VIGAS VI TAA IOHHA GARAGA SAA
X59764; (X66133)	
	(APEN) (REF-1 PROTEIN) (APE 1)
079718	ENDONUCLEASE III HOMOLOG 1 (HNTH1) (OCTS3)
7404514	DNA LIGASE I (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) (DNL 1) (LIG 1)
X84740	DNA LIGASE III (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) (DNL3)
M18112	POLY (ADP-RIBOSE) POLYMERASE (PARP) (ADPRI) (NAD (+) ADP-RIBOSYL IIKANSFEKASE) (POLY (ADP-RIBOSE) SYNTHETASE) (PPOL)
D16581	7,8-DIHYDRO-8-OXOGUANINE TRIPHOSPHATASE (mutt HOMOMOLOG) (8-OXO-IOGTPASE) (MIH1)
M36089	DNA-REPAIR PROTEIN XRCC1
D29013	DNA POLYMERASE BETA (DPOB)
M11722	DNA NUCLEOTIDYLEXOTRANSFERASE (TERMINAL ADDITION ENZYME) (TERMINAL DEOXYNUCLEOTIDYLTRANSFERASE) (TERMINAL TRANSFERASE) (DNIT) (TDT)
X55715	40S RIBOSOMAL PROTEIN S3 (POSSIBLE dispase)
	NUCLEO II DE EXCISION REPAIX

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
D14533	DNA-REPAIR PROTEIN COMPLEMENTING XP-A CELLS (XERODERMA PIGMENTOSUM GROUP A COMPLEMENTING PROTEIN)
M31899	DNA-REPAIR PROTEIN COMPLEMENTING XP-B CELLS (XERODERMA PIGMENTOSUM GROUP B COMPLEMENTING PROTEIN) (DNA EXCISION REPAIR PROTEIN ERCC3) (BASAL TRANSCRIPTION FACTOR 2 89 KD SUBUNIT) (BTF2-p89) (TFIIH 89 KD SUBUNIT)
D21089	DNA-REPAIR PROTEIN COMPLEMENTING XP-C CELLS (XERODERMA PIGMENTOSUM GROUP C COMPLEMENTING PROTEIN) (p.125)
D21235 D21090	UV EXCISION REPAIR PROTEIN PROTEIN RAD23 HOMOLOG A (HHR23A) UV EXCISION REPAIR PROTEIN PROTEIN RAD23 HOMOLOG B (HHR23B) (XP-C REPAIR COMPLEMENTING COMPLEX 58 KD PROTEIN) (p58)
X52221; (HT1175)	DNA-REPAIR PROTEIN COMPLEMENTING XP-D CELLS (XERODERMA PIGMENTOSUM GROUP D COMPLEMENTING PROTEIN) (DNA EXCISION REPAIR PROTEIN ERCC-2)
U18299	DAMAGE-SPECIFIC DNA BINDING PROTEIN p.127 SUBUNIT; IMPLICATED IN XERODERMA PIGMENTOSUM GROUP E (DDB1)
018300	DAMAGE-SPECIFIC DNA BINDING PROTEIN p48 SUBUNIT; IMPLICATED IN XERODERMA PIGMENTOSUM GROUP E (DDB2)
177890	DNA-REPAIR PROTEIN COMPLEMENTING XP-F CELLS (XERODERMA PIGMENTOSUM GROUP F COMPLEMENTING PROTEIN) (DNA EXCISION REPAIR PROTEIN ERCC-4)
L20046; (X69978)	DNA-REPAIR PROTEIN COMPLEMENTING XP-G CELLS (XERODERMA PIGMENTOSUM GROUP G COMPLEMENTING PROTEIN) (DNA EXCISION REPAIR PROTEIN ERCC-5)
U28413	COCKAYNE SYNDROME GROUP A: WD-REPEAT PROTEIN (CSA PROTEIN)
104791	EXCISION REPAIR PROTEIN EXCC-6 (C3B) BASIC TRANSCRIPTION FACTOR 42 KD STRINIT (PAS) (RTF20A2)
230094	BASIC TRANSCRIPTION FACTOR 2, 44 KD SUBUNIT (BIF2p44)
230093	BASIC TRANSCRIPTION FACTOR 2, 34 KD SUBUNIT (BIF2p34)
Y0/595 M13194	BASIC HANSCRIPTION FACTOR 2, 32 RD SUBUNIT (BIT-2032) DNA EXCISION REPAIR PROTEIN ERCC-1

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
M63488	REPLICATION PROTEIN A 70 KD DNA-BINDING SUBUNIT (RP-A) (REPLICATION FACTOR-A PROTEIN 1) (SINGLE STRANDED DNA-BINDING PROTEIN)
305249	REPLICATION PROTEIN A 32 KD SUBUNIT (RP-A) (RF-A) (REPLICATION FACTOR-A PROTEIN 2)
107493	REPLICATION PROTEIN A 14 KD SUBUNIT (RP-A) (RF-A) (REPLICATION FACTOR A PROTEIN 3)
U24186	REPLICATION PROTEIN A 30 KD SUBUNIT (RP-A) (RF-A) (REPLICATION FACTOR-A PROTEIN A)
; (J04718)	ROLIFERATING
107540	ACTIVATOR 1 36 KD SUBUNIT (REPLICATION FACTOR C 36 KD SUBUNIT) (RFC36) ACTIVATOR 1 37 KD SUBUNIT (REPLICATION FACTOR C 37 KD SUBUNIT) (RFC37)
	1
M87338	ΙI
L14922	ACTIVATOR 1 140KD SUBUNIT (REPLICATION FACTOR C LARGE SUBUNIT) (A1 140 KD SUBUNIT) (RF-C 140 KD SUBUNIT) (ACTIVATOR 1 LARGE SUBUNIT) (DNA-BINDING PROTEIN PO-GA)
X06745	DNA POLYMERASE ALPHA
M80397	DNA POLYMERASE DELTA CATALYTIC CHAIN
M60974	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD-45 (DNA-DAMAGE INDUCIBLE TRANSCRIPT 1) (DDIT1) (GA45)
\$40706 (\$62138)	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD 153 (DNA-DAMAGE INDITIONS F PROTEIN) (CHOP).
	Homologous recombination
U63139	DNA REPAIR PROTEIN RAD50
D13804: (D14134)	D13804: (D14134) DNA REPAIR PROTEIN RAD51 HOMOLOG
U12134	DNA REPAIR PROTEIN RAD52 HONOLOG
U09820	X-LINKED HELICASE II (X-LINKED NUCLEAR PROTEIN) (XNP) (RADS4L) (XH2)
x97795	DNA REPAIR PROTEIN RADS4 HOMOLOG
U14680	BREAST CANCER TYPE 1 SUSCEPTIBILITY PROTEIN (BRCA1)
U43746	BREAST CANCER TYPE 2 SUSCEPTIBILITY PROTEIN (BRCA2)
D63882	MEIOTIC RECOMBINATION PROTEIN DMC1/LIM15 HOMOLOG
X83441	DNA LIGASE IV (POLYDEOXYRIBONUCLEOTIDE SYNTHASE (ATP)) (DNL4)

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
M74524	HHR6A (YEAST RAD6 HOMOLOG) (UBIQITIN-CONJUGATING ENZYME) (UBCA)
	HHRAB (YEAST RADA HOMOLOG) (UBIQITIN-CONJUGATING ENZYME) (UBCB)
Y08837	RADS1-LIKE PROTEIN (POSSIBLE XRCC2)
	Non-homologous end-rejoining
U40622	DNA REPAIR PROTEIN XRCC4
M32865; (S38729)	ATP-DEPENDENT DNA HELICASE II, 70 KD SUBUNIT (LUPUS KU AUTOANTIGEN PROIEIN P70) (70 KD SUBUNIT OF KU ANTIGEN) (THYROID-LUPUS AUTO-ANTIGEN) (TLAA) (KU70)
	(CTC BOX BINDING FACTOR 75 KD SUBUNIT) (CTCBF) (CTC75) (XRCC6)
M30938	ATP-DEPENDENT DNA HELICASE IL 86 KD SUBUNIT (LUPUS KU AUTOANTIGEN PROTEIN
	P86) (86 KD SUBUNIT OF KU ANTIGEN) (THYROID-LUPUS AUTOANTIGEN) (TLAA) (CIC BOX) BINDING FACTOR 85 KD SUBUNIT) (CTCBF) (CTCBS) (NUCLEAR FACTOR IV) (KUBO) (XRCC5)
(7020)	DNA-DEPENDENT PROTEIN KINASE (DNA-PK)
	DNA DEPENDENT PROTEIN KINASE CATALYTIC SUBUNIT (DNA-PKCS) (XRCC7)
M29474	V(D)J RECOMBINATION ACTIVATING PROTEIN 1 (RAG1) (RAG-1)
M94633	V(D)J RECOMBINATION ACTIVATING PROTEIN 2 (RAG2) (RAG-2)
	MISMATCH REPAIR
U07418; (U07343)	DNA MISMATCH REPAIR PROTEIN MILH (mutl. HOMOLOG)
UD4045; (L47583)	DNA MISMATCH REPAIR PROTEIN MSH2
J04810	DNA MISMAICH KEPAIK PKOIEIN MATA (DIVEKGEN) OF SINCANT KOLE. 9 (MISMAICH PROIEIN 1) (MRP1) (DUG)
U54777	DNA MISMATCH REPAIR PROTEIN MSH6 (muts - ALPHA 160 KD SUBUNIT) (G/f MISMATCH
113404	BINDING PROTEIN (GIBP) (GIMBP) (FIDS) DNA MISMATCH REPAIR PROTEIN PMS2 (PMS1 PROTEIN HOMOLOG 2)
2605	
U13695	DNA MISMATCH REPAIR PROTEIN PMS1 (PMS1 PROTEIN HOMOLOG 1)
X14672; X17059	ARYLAMINE N-ACETYLTRANSFERASE, POLYMORPHIC (EC. 2.3.1.3) (PNA1) + ARYLAMINE N-ACETYLTRANSFERASE, MONOMORPHIC (EC. 2.3.1.5) (MNA1)
200036	CYTOCHROME PA50 IA2 (EC 1.14.14.1) (P450-P3) (P450-4).515
200036	CYTOCHROME P450 IA2 (EC 1.14.14.1) (P450-P3) (P450-4).515

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
J04449; D00003; J04813; D00408	CYTOCHROME PA50 IIIAA (EC 1.14.14.1) (NIFEDIPINE OXIDASE) (NF-25) (PA50-PCN1)
	CYTOCHROME PA50 IIIA3 (EC 1.14.14.1) (GLUCOCORTICOID-INDUCIBLE) (HLP) CYP3A3.
	CYTOCHROME P450 IIIA5 (EC 1.14.14.1) (P450-PCN3)
	CYTOCHROME PA50 IIIA7 (EC 1.14.14.1) (PA50-HFLA)
102871	CYTOCHROME P450 IVB1 (EC 1.14.14.1) (P450-HP)
M33318; (X13930;	CYTOCHROME P450 IIA6 (EC 1.14.14.1) (COUMARIN 7-HYDROXYLASE) (IIA3) (P450(I)) (PHENORARRIAL-INDLICIRE)
7	CYTOCHROME P450 IIA7 (EC 1.14.14.1) (P450-IIA4)
P450 IIA7 (EC	
1.14.14.1) (14.05	
M21940; M15331;	CYTOCHROME P450 IIC9 (EC 1.14.14.1) (P450 PB-1) (P450 MP-4) (S-MEPHENYTOIN 4-
(M21939)M61858; HYDROXYLASE)	HYDROXYLASE)
(L07093); M61853; M61854	(L07093); M61853; CYTOCHROME P450 !} M61854
921600	DIHYDROPYRINIDINE DEHYDROGENASE (NADP+) PRECURSOR (EC 1.3.1.2) (DPD) (DIHYDROURACIL DEHYDROGENASE) (DIHYDROTHYMINE DEHYDROGENASE) DPYD.
	INTERNATION OF LATER CONTRACT TO THE PROPERTY OF THE PROPERTY
M64082	DIMETHYLANIUNE MONOCATGERASE (N-CAIDE FORMING) 1 (EC. 1.14.13.6) (TELAL HEPATIC FLAVIN-CONTAINING MONOCAYGENASE 1) (FMO 1) (DIMETHYLANIUNE OXIDASE 1)
M83772	DIMETHYLANIUNE MONOOXYGENASE (N-OXIDE FORMING) 3 (EC 1.14.13.8) (HEPATIC
	FLAVIN-CONTAINING MONOOXYGENASE 3) (FMO 3) (DIMETHYLANILINE OXIDASE 3) (FMO II)
211737	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 4 (EC 1.14.13.8) (HEPATIC
	FLAVIN-CONTAINING MONOOXYGENASE 4) (FMO 4) (DIMETHYLANILINE OXIDASE 4)
137080	DIMETHYLANILINE MONOOXYGENASE (N-OXIDE FORMING) 5 (EC 1.14.13.8) (HEPATIC IS A VIN-CONTAINING MONOOXYGENASE 5) (FMO 5) (DIMETHYLANILINE OXIDASE 5)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
X04808	PORPHOBILINOGEN DEAMINASE (EC 4.3.1.8) (HYDROXYMETHYLBILANE SYNTHASE) (HMBS) (PRE-UROPORPHYRINOGEN SYNTHASE)
M14758	MULTIDRUG RESISTANCE PROTEIN 1 (P-GLYCOPROTEIN 1)
M23234	MULTIDRUG RESISTANCE PROTEIN 3 (P-GLYCOPROTEIN 3)
105628	MULTIDRUG RESISTANCE-ASSOCIATED PROTEIN 1
1208021	NICOTINAMIDE N-METHYLTRANSFERASE (EC 2.1.1.1)
U09031; U28170; L19956	PHENOL-SULFATING PHENOL SULFOTRANSFERASE 1 (EC 2.8.2.1) (P-PSI) (THERMOSTABLE PHENOL SULFOTRANSFERASE) (TS-PST) (HAST1/HAST2) (ST1A3) STP1 OR STP.
	PHENOL-SULFATING PHENOL SULFOTRANSFERASE 2 (EC 2.8.2.1) (P-PST) (ST1A2) STP2.
	MONOAMINE-SULFATING PHENOL SULFOTRANSFERASE (EC 2.8.2.1) (SULFOTRANSFERASE, MONOAMINE-PREFERRING) (M-PST) (THERMOLABILE PHENOL SULFOTRANSFERASE) (TL-PST) (PLACENTAL ESTROGEN SULFOTRANSFERASE) (CATECHOLAMINE-SULFATING PHENOL SULFOTRANSFERASE) (HAST3) STM.
U08854; X63359; U06641; J05428; Y00317	UDP-GLUCURONOSYLTRANSFERASE 2815 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGTH-3) UGT2815.
	UDP-GLUCURONOSYLIRANSFERASE 2810 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) UGT2810.
	UDP-GLUCURONOSYLTRANSFERASE 288 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (ESTRIOL SPECIFIC) (HLUG4) (FRAGMENT) UGT288.
	UDP-GLUCURONOSYLTRANSFERASE 287 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (3.4-CATECHOL ESTROGEN SPECIFIC) (UDPGTH-2) UGT287.
	UDP-GLUCURONOSYLTRANSFERASE 284 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (HYODEOXYCHOUC ACID) (HLUG25) (UDPGTH-1) UGT284.

TABLE 5 (CONT)

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
M68840	AMINE OXIDASE (FLAVIN-CONTAINING) A (EC 1.4.3.4) (MONOAMINE OXIDASE) (MAO-
M69177	A) MACJA. AMINE OXIDASE (FLAVIN-CONTAINING) B (EC 1.4.3.4) (MONOAMINE OXIDASE) (MAO- B) MAOB.
K03191	CYTOCHROME P450 IA1 (EC 1.14.14.1) (P450-P1) (P450 FORM 6) (P450-C) (TCDD-INDUCIBLE).
M29874	CYTOCHROME P450 IIB6 (EC 1.14.14.1) (PHENOBARBITAL-INDUCIBLE) (P450 IIB1).
	CYTOCHROME P450 IID6 (EC 1.14.14.1) (P450-DB1) (DEBRISOQUINE 4-HYDROXYLASE) CYP2D6.
J02625	CYTOCHROME P450 IIE1 (EC 1.14.14.1) (P450-J) (ETHANOL INDUCIBLE) CYP2E1
202906	CYTOCHROME P450 IIF1 (EC 1,14.14.1) CYP2F1.
M14565	CYTOCHROME P450 XIA1, MITOCHONDRIAL PRECURSOR (EC 1.14.15.6) (P450(SCC)) (CHOLESTEROL SIDE-CHAIN CLEAVAGE ENZYME) (CHOLESTEROL DESMOLASE)
x55764	CYTOCHROME P450 XIB1 PRECURSOR (P450C11) (STEROID 11-BETA-HYDROXYLASE) (EC11,14,15,4) CYP11B1 OR S11BH.
M12792; (M23280)	M12792; (M23280) CYTOCHROME P450 XXIB (EC 1.14.99.10) (STEROID 21-HYDROXYLASE) (P450-C21B) CYP21B OR CYP21 OR CYP21A2.
107765	LIVER CARBOXYLESTERASE PRECURSOR (EC 3.1.1.1) (ACYL COENZYME A:CHOLESTEROL ACYLTRANSFERASE) (ACAT) (MONOCYTE/MACROPHAGE SERINE ESTERASE) (HMSE)
J05459	GLUIATHIONE S-TRANSFERASE MU 3 (EC 2.5.1.18) (GSTM3-3) (CLASS-MU) GSTM3 OR GSTS.
D13889	GLUTATHIONE REDUCTASE
X15722	욁
J03746	GLUTATHIONE S-TRANSFERASE M4 (GLUTATHIONE S-TRANSFERASE MU 1)
X08020	GLUTATHIONE S-TRANSFERASE P CLUTATHIONE S-TRANSFERASE A1-1 (Glutothione S-transferase (GST) Ha subunit 1)
M14777	1
M21304	GLUTHATHIONE S-TRANSFERASE (THETA 1) CLITATHIONE-S-TRANSFERASE HOMOLOG
6/2907	SOLUBLE EPOZIDE HYDROLASE (SEH) (EC 3.3.2.3) (EPOXIDE HYDRATASE) (CYTOSOLIC
	EPONDE NIDROLLASS (CELL) EL 1965.

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
M57899	UDP-GLUCURONOSYLÍRANSFERASE 1-1 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (UGT-1A) (UGT1-1) (UGT1-01) (UGT1.1) (UGT1A1) (BIURUBIN SPECIFIC ISOZYME 1) (UGT1A) (HUG-BR1) UGT1 OR GNT1.
S55985	UDP-GLUCURONOSYLTRANSFERASE 1-2 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPG.), (UGT-1B) (UGT1-2) (UGT1-02) (UGT1.2) (UGT1A2) (UGT1B) (HLUGP4) UGT1 OR GNT1.
M84127	UDP-GLUCURONOSYLTRANSFERASE 1-3 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGT) (UGT-1C) (UGT1-3) (UGT1-03) (UGT1A3) (UGT1C) UGT1 OR GNT1.
M57951	UDP-GLUCURONOSYLTRANSFERASE 1-4 PRECUISOR, MICROSOMAL (EC 2.4.1.17) (UDPGI) (UGT-1D) (UGT1-4) (UGT1-04) (UGT1.4) (UGT1A4) (UGT1D) (BILIRUBIN SPECIFIC ISOZYME 2) (HUG-BR2) UGT1 OR GNT1.
J040 9 3	UDP-GLUCURONOSYLTRANSFERASE 1-6 PRECURSOR, MICROSOMAL (EC 2.4.1.17) (UDPGI) (UGT-1F) (UGT1-6) (UGT1-6) (UGT1-6) (UGT1-6) (UGT1-6) (UGT1-7)
X71480	CYTOCHROME P450 IVA11 (EC 1.14.14.1) (FRAGMENT) CYP4A-11.
X83573	ARYLSULFATASE E PRECURSOR (EC 3.1.6) (ASE) ARSE.
X92106	BLEOMYCIN HYDROLASE (EC 3.4.22) (BLM HYDROLASE).
M65212	CATECHOL O-METHYLTRANSFERASE, MEMBRANE-BOUND FORM (EC 2.1.1.6) (MB-COMT) (CONTAINS: CATECHOL O-METHYLTRANSFERASE, SOLUBLE FORM (S-COMT)) COMT.
228409	COPROPORPHYRINOGEN III OXIDASE PRECURSOR (EC 1.3.3.3) (COPROPORPHYRINOGENASE) (COPROGEN OXIDASE) (COX) CPO.
Y09501	NADH-CYTOCHROME B5 REDUCTASE (EC 1.6.2.2) (B5R) DIA1.
U12778	ACYL-COA DEHYDROGENASE, SHORT/BRANCHED CHAIN SPECIFIC PRECURSOR (EC 1.3.99) (SBCAD) (2-METHYL BRANCHED CHAIN ACYL-COA DEHYDROGENASE) (2-MEBCAD) ACADSB.
M74542	ALDEHYDE DEHYDROGENASE, DIMERIC NADP-PREFERRING (EC 1.2.1.5) (CLASS 3). ALDH3.

GenBank #	STRESS RESPONSE REGULATORS AND EFFECTORS
X53463	GLUTATHIONE PEROXIDASE-GASTROINTESTINAL (EC 1.11.1.9) (GSHPX-GI) (GLUTATHIONE PEROXIDASE-RELATED PROTEIN 2) (GPRP) GPX2.
X71973	PHOSPHOLIPID HYDROPEROXIDE GLUTHATIONE PEROXIDASE (EC 1.11.1.9) (PHGPX) GPX4.
M63012	SERUM PARAOXONASE/ARYLESTERASE 1 (EC 3.1.1.2) (EC 3.1.8.1) (PON 1) (SERUM ARYLDIAKYLPHOSPHATASE 1) (A-ESTERASE 1) (AROMATIC ESTERASE 1) PON 1 OR PON.
148513	SERUM PARAOXONASE/ARYLESTERASE 2 (EC 3.1.1.2) (EC 3.1.8.1) (PON 2) (SERUM ARYLDIAKYLPHOSPHATASE 2) (A-ESTERASE 2) (AROMATIC ESTERASE 2) PON2.
148516	SERUM PARAOXONASE/ARYLESTERASE 3 (EC 3.1.1.2) (EC 3.1.8.1) (PON 3) (SERUM ARYLDIAKYLPHOSPHATASE 3) (A-ESTERASE 3) (AROMATIC ESTERASE 3) (FRAGMENT) PON3.
262904	THIOPURINE S-METHYLTRANSFERASE (EC 2.1.1.67) (THIOPURINE METHYLIRANSFERASE) TPMI.
102932	PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR ALPHA (PPAR-ALPHA) PPARA OR PPAR
107592	PEROXISOME PROLIFERATOR ACTIVATED RECEPTOR BETA (PPAR-BETA) (NUCLEAR HORMONE RECEPTOR 1) (NUC1) (NUC1) PPARB OR PPARD.
	HOUSEKEEPING GENES
M26880	UBIQUITIN DAICCHOLI IDACE A2
V00530	HYPOXANIHINE-GUANINE PHOSPHORIBOSYLTRANSFERASE
X01677	GLYCERALDEHYDE 3-PHOSPHATE DEHYDROGENASE
K00558	TUBULIN ALPHA
M11886	HLA CLASSI HISIOCOMPAIIBILITY ANTIGEN, C-4 ALTRA CRAIN
(MHC) X00351	BETA-ACTIN
X56932	23 KD HIGHLY BASIC PROTEIN
U14971	RIBOSOMAL PROTEIN S9
	NEGATIVE CONTROLS
	INCOMINE COMINCIE

WO 98/53103 PCT/US98/10561

Oncogene and Tumor Suppressor Gene Array

In the oncogene and tumor suppressor gene array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that are associated with cellular proliferative diseases, specifically neoplastic diseases. Genes of interest that may be represented on the array include: oncogenes and tumor suppressor genes. In a specific oncogene and tumor suppressor gene array of interest, the spots are as provided in Table 6.

TABLE 6

MYC PROTO-ONCOGENE PROTEIN HER3 (ERB-B3)[Epidermal growth factor b) oncogene homolog)] INSULIN-LIKE GROWTH FACTOR I RE MACROPHAGE COLONY STIMULATIR (PDGF-2) (BACAPLERMIN) (C-SIS) THANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN EAR-1 1) (X00588; EPIDERMAL GROWTH FACTOR REC COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 (EPIDERMAL GROWTH FA THANSCRIPTION FACTOR RECEFTOR PROTEIN FA TUMOR NECROSIS FACTOR RECEFTOR PROTEIN PROTEIN FA TUMOR NECROSIS FACTOR PROTEIN FA TUMOR NECROSIS FACTOR PROTEIN FA			Contains
M2936	Cendant		Gelle Name
M29366 HER3 (ERB-B3)[Epidemal growth factor	V00568		MYC PROTO-ONCOGENE PHOLEIN
X04434 INSULIN-LIKE GROWTH FACTOR I RE X03663 MACROPHAGE COLONY STIMULATII X02811; [X02744; PLATELET-DERIVED GROWTH FACTOR U33201) X02811; [X02744; PLATELET-DERIVED GROWTH FACTOR TNF X01394 TOMOR NECROSIS FACTOR TNF X01394 TRANSFORMING GROWTH FACTOR TNF X02812 TRANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR X02812 TRANSFORMING GROWTH FACTOR X02812 TRANSFORMING GROWTH FACTOR X02812 TRANSFORMING GROWTH FACTOR X02812 TG-RETA RECEPTOR TYPE III X07282 TGF-BETA RECEPTOR TYPE III X07282 TGF-BETA RECEPTOR RETA-2 X07282 TGF-BETA RECEPTOR RETA-2 X07282 TGF-BETA RECEPTOR TYPE III X07282 TGF-BETA RECEPTOR TYPE III X07282 TGF-BETA RECEPTOR TYPE III X07283 TGF-BETA RECEPTOR TOTAL RECASES X12794 TGF-BBA RELATED PROTEIN EAR-2 X12794 COUP TRANSCRIPTION FACTOR U11732 EFBBA RELATED PROTEIN TEL U11732 ETS-RELATED PROTEIN TEL U11732 ETS-RELATED PROTEIN TEL U11734 ETS-RELATED PROTEIN TEL U11735 ETS-RELATED PROTEIN TEL U11730 E	M29366		HER3 (ERB-B3)[Epidermal growth factor receptor (avian erythroblastic leukemia viral (v-erb-b) oncogene homolog)]
MACROPHAGE COLONY STIMULATING			INSULIN-LIKE GROWTH FACTOR I RECEPTOR
212020; [M92424] MDM2 PROTEIN (PS3-ASSOCIATED PROTEIN) + MDM2-A (GB: U33201) M12783 PLATELET-DERIVED GROWTH FACTOR, B CHAIN PRECURSO M127894 TUMOR-Q: [BACAPLERMIN] (C-SIS) X02812 TRANSFORMING GROWTH FACTOR BETA [1] M15024 MYB PROTO-ONCOGENE PROTEIN M15024 MYB PROTO-ONCOGENE PROTEIN M15024 MYB PROTO-ONCOGENE PROTEIN M15024 TRANSFORMING GROWTH FACTOR BETA [1] M15024 TRANSFORMING GROWTH FACTOR BETA [1] M15024 TRANSFORMING GROWTH FACTOR BETA [2] X06182 C-kit TRANSFORMING GROWTH FACTOR BETA [2] X06182 C-kit TRANSFORMING GROWTH FACTOR BETA [2] X06182 TGF-BETA RECEPTOR TYPE III X06182 C-kit TGF-BETA RECEPTOR PROTEIN BETA-2 X13293 MYB-RELATED PROTEIN BETA-2 X13293 W-ERBA RELATED PROTEIN BERA-2 TCHABA RELATED PROTEIN EAR-1 TCHYroid hormone trilodoth V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR RECEPTOR PRECEPTOR TO-SEBA TRANSCRIPTION FACTOR RECEPTOR TO-SEBA TRANSCRIPTION FACTOR RECEPTOR TO-SEBA TRANSCRIPTION FACTOR RECEPTOR TO-SEBA TRANSCRIPTION FACTOR RECEPTOR TUBASCRIPTION FACTOR RECEPTOR TRANSCRIPTION FACTOR RECEPTOR TUBASCRIPTION FACTOR RECEPTOR TUBASCRIPTION FACTOR RECEPTOR TRANSCRIPTION FACTOR RECEPTOR TUBASCRIPTION FACTOR TUBASCRIPTION FACTOR	X03663		MACROPHAGE COLONY STIMULATING FACTOR I RECEPTOR [c-fms proto-oncogene]
X02811; [X02744; PLATELET-DERIVED GROWTH FACTOR, B CHAIN PRECURSO] M12783] (PDGF-2) (BACAPLERMIN) (C-SIS) X01394 TUMOR NECROSIS FACTOR [TNFa] X02812 TRANSFORMING GROWTH FACTOR BETA [1] M15024 MYB PROTO-ONCOGENE PROTEIN M14594 CELLULAR TUMOR ANTIGEN PS3 M19154 TRANSFORMING GROWTH FACTOR BETA [2] X07282 RETINOIC ACID RECEPTOR TYPE III X07282 RETINOIC ACID RECEPTOR BETA-2 X13293 WYB-RELATED PROTEIN EAR-1 [Thyroid hormone triodoth) M24698 V-ERBA RELATED PROTEIN EAR-1 [Thyroid hormone triodoth) X12794 COUP TRANSCRIPTION FACTOR RECEPTOR PRECURSOR (EC X12795 COUP TRANSCRIPTION FACTOR RECEPTOR [V-rbA related ear-3 protein) U11732 ETS-RELATED PROTEIN TEL U1782 ETS-RELATED PROTEIN TEL U17842 DP2 (Humdp2), dimerization partner of E2F LO7668 ERBB4 [EPIDERMAL GROWTH FACTOR RECEPTOR] M33294 TUMOR NECROSIS FACTOR RECEPTOR M41730 ERBB-2 RECEPTOR PROTEIN-TROSINE KINASE	Z12020;	[M92424]	MDM2 PROTEIN (PS3-ASSOCIATED PROTEIN) + MDM2-A (GB: U33199) + MDM2-C (GB: U33201)
THANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR MYB PROTO-ONCOGENE PROTEIN CELLULAR TUMOR ANTIGEN P53 TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR B [8-myb] V-ERBA RELATED PROTEIN EAR-1 1] (X00588; EPIDERMAL GROWTH FACTOR REC U48722] (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 [EPIDERMAL GROWTH FA TRANSCRIPTION FACTOR AP-1 [6- TUMOR NECROSIS FACTOR RECEF FRBB-2 RECEPTOR PROTEIN-TYRC	X02811;	[X02744;	PLATELET-DERIVED GROWTH FACTOR, B CHAIN PRECURSOR (PDGF B-CHAIN)
TRANSFORMING GROWTH FACTOR TRANSFORMING GROWTH FACTOR MYB PROTO-ONCOGENE PROTEIN CELLULAR TUMOR ANTIGEN P53 TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR BETA-2 MYB-RELATED PROTEIN B [B-myb] V-ERBA RELATED PROTEIN EAR-1 1) (EABB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERB4 [EPIDERMAL GROWTH FA THANSCRIPTION FACTOR RECEPTOR BEGET TUMOR NECROSIS FACTOR RECEFTOR PROTEIN TEL TUMOR NECROSIS FACTOR RECEFTOR PROTEIN-TYRC	X01394		TUMOR NECROSIS FACTOR [TNFa]
TRANSFORMING GROWTH FACTOR MYB PROTO-ONCOGENE PROTEIN CELLULAR TUMOR ANTIGEN P53 TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B [B-myb] V-ERBA RELATED PROTEIN EAR-1 1) [X00588; EPIDERMAL GROWTH FACTOR REC U48722] (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 [EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 [c- TUMOR NECROSIS FACTOR RECEF	K03222		TRANSFORMING GROWTH FACTOR-ALPHA
MYB PROTO-ONCOGENE PROTEIN CELLULAR TUMOR ANTIGEN P53 TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B (B-myb) V-ERBA RELATED PROTEIN EAR-1 1) (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner o ERBB4 [EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 fc- TUMOR NECROSIS FACTOR RECEF	X02812		TRANSFORMING GROWTH FACTOR BETA [1]
CELLULAR TUMOR ANTIGEN P53 TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B (B-myb) V-ERBA RELATED PROTEIN EAR-1 1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 (EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 (c- TUMOR NECROSIS FACTOR RECEF FRBB-2 RECEPTOR PROTEIN-TYRC	M15024		MYB PROTO-ONCOGENE PROTEIN
TRANSFORMING GROWTH FACTOR C-kit TGF-BETA RECEPTOR TYPE III TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B (B-myb) V-ERBA RELATED PROTEIN EAR-1 1) (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 (EPIDERMAL GROWTH FA TRANSCRIPTION FACTOR AP-1 (c- TUMOR NECROSIS FACTOR RECEF FRBB-2 RECEPTOR PROTEIN-TYRC	M14694		CELLULAR TUMOR ANTIGEN P53
C-kit TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B [B-myb] V-ERBA RELATED PROTEIN EAR-1 1) (X00588; EPIDERMAL GROWTH FACTOR REC U48722] (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner o ERBB4 [EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 [c-	M19154		TRANSFORMING GROWTH FACTOR BETA [2]
TGF-BETA RECEPTOR TYPE III RETINOIC ACID RECEPTOR BETA-2 MYB-RELATED PROTEIN B (B-myb) V-ERBA RELATED PROTEIN EAR-1 1) (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner o ERBB4 (EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 (c-	X06182		Q-Kit
MYB-RELATED PROTEIN B [8-myb] V-ERBA RELATED PROTEIN B [8-myb] V-ERBA RELATED PROTEIN EAR-1 (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 [EPIDERMAL GROWTH FA THANSCRIPTION FACTOR AP-1 [6-10] TUMOR NECROSIS FACTOR RECEFFICATION PROTEIN TECEFFICATION FACTOR RECEFFICATION FACTOR RECEFFICATION PROTEIN-TYRC	107594		TGF-BETA RECEPTOR TYPE III
WYB-RELATED PROTEIN B (8-myb) V-ERBA RELATED PROTEIN EAR-1 1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 (EPIDERMAL GROWTH FATOR PROTEIN TEL TRANSCRIPTION FACTOR AP-1 (c- TUMOR NECROSIS FACTOR RECEF	X07282		RETINOIC ACID RECEPTOR BETA-2
(X00588; EPIDERMAL GROWTH FACTOR REC U48722] (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of ERBB4 [EPIDERMAL GROWTH FA TRANSCRIPTION FACTOR AP-1 [6- TUMOR NECROSIS FACTOR RECEF	X13293		MYB-RELATED PROTEIN B (B-myb)
VERBA RELATED PROTEIN EAR-2 V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR [V-erbA RETS-RELATED PROTEIN TEL	M24898		V-ERBA RELATED PROTEIN EAR-1 [Thyroid hormone triiodothyronine receptor c-erbA,ear-
U48722) (ERBB1) V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR [V-erbA r ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of E2F ERBB4 [EPIDERMAL GROWTH FACTOR R TRANSCRIPTION FACTOR AP-1 [c-jun proto TUMOR NECROSIS FACTOR RECEPTOR	700400	. IVOCEBB.	1) EDINEBMAL GROWTH FACTOR RECEPTOR PRECIIRSOR (FC 2.7.1.12). (FGFR)
V-ERBA RELATED PROTEIN EAR-2 COUP TRANSCRIPTION FACTOR [V-erba raction of the county	X00663;	; [AW306, ; U48722]	(ERBB1)
COUP TRANSCRIPTION FACTOR [V-erbA1] ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of E2F ERBB4 [EPIDERMAL GROWTH FACTOR R TRANSCRIPTION FACTOR AP-1 [c-jun proto TUMOR NECROSIS FACTOR RECEPTOR	X12794		V-ERBA RELATED PROTEIN EAR-2
ETS-RELATED PROTEIN TEL DP2 (Humdp2), dimerization partner of E2F ERBB4 [EPIDERMAL GROWTH FACTOR R THANSCRIPTION FACTOR AP-1 [c-jun proto TUMOR NECROSIS FACTOR RECEPTOR	X12795		COUP TRANSCRIPTION FACTOR [V-erbA related ear-3 protein]
DP2 (Humdp2), dimerization partner of E2F ERBB4 [EPIDERMAL GROWTH FACTOR R THANSCRIPTION FACTOR AP-1 [c-jun proto TUMOR NECROSIS FACTOR RECEPTOR ERBB-2 RECEPTOR PROTEIN-TYROSINE KII	U11732		ETS-RELATED PROTEIN TEL
TRANSCRIPTION FACTOR AP-1 (c-jun proto TUMOR NECROSIS FACTOR RECEPTOR FERBE-2 RECEPTOR PROTEIN-TYROSINE KII	U18422		DP2 (Humdp2), dimerization partner of E2F
THANSCRIPTION FACTOR AP-1 (c-jun proto TUMOR NECROSIS FACTOR RECEPTOR FEBB-2 RECEPTOR PROTEIN-TYROSINE KII	L07868		1
TUMOR NECROSIS FACTOR RECEPTOR FRBB-2 RECEPTOR PROTEIN-TYROSINE KII	J04111		TRANSCRIPTION FACTOR AP-1 [c-jun proto oncogene]
	M33294		TUMOR NECROSIS FACTOR RECEPTOR [Tumor necrosis factor receptor 1 (55kD)]
	M11730	0	ERBB-2 RECEPTOR PROTEIN-TYROSINE KINASE

TABLE 6 (CONT)

L12260	HEREGULIN ALPHA [Recombinant glial growth factor 2]
L12261	HEREGULIN ALPHA [Recombinant glial growth factor]
M27288	ONCOSTATIN M
M59964	STEM CELL FACTOR (C-KIT LIGAND)
M76125	AXL (TYROSINE-PROTEIN KINASE RECEPTOR UFO)
X06182	C-KIT PROTO-ONCOGENE [mast/stem cell growth factor receptor]
X06374	PLATELET-DERIVED GROWTH FACTOR A CHAIN
D13866	ALPHA-CATENIN
D17517	SKY (DTK) (TYRO3) (RSE)
L11353; Z22664;	MERLIN (SCHWANNOMIN) (moesin-ezrin-radixin-ilike protein)(neurotibromatosis 2)
X72657; L27133	١
L13738	TYROSINE-PROTEIN KINASE SYK activated p21cdc42rls killase (ach))
L14837	- 1
L16785	NUCLEOSIDE DIPHOSPHATE KINASE B [c-myc transcription factor (put)]
L19067	TRANSCRIPTION FACTOR P65
120422	PROTEIN ETA (14-3-3 PROTEIN ETA)
122075	GUANINE NUCLEOTIDE REGULATORY PROTEIN (G13)
125259	T LYMPHOCYTE ACTIVATION ANTIGEN CD86 [CD28 antigen ligand 2, B7-2 antigen]
1.33264	CDC2-RELATED KINASE PISSLRE
M13150	MAS PROTO-ONCOGENE
M31213; [M5746	M31213; IM57464] PROTO-ONCOGENE TYROSINE-PROTEIN KINASE RECEPTOR RET PRECURSOR (EC
M31899	DNA-REPAIR PROTEIN COMPLEMENTING XP-B CELLS [DNA repair helicase (ERCC3)]
M32865	ATP-DEPENDENT DNA HELICASE II (70 KD SUBUNIT) [Thyroid autoantigen 70kD (Ku
	antigen))
M34960	TRANSCRIPTION FACTOR IID
M36089	DNA-REPAIR PROTEIN XRCC1
M54915	PIM-1 PROTO-ONCOGENE (SERINE/THREONINE-PHOLEIN KINASE)
M60915	NEUROFIBROMIN [neurofibromatosis protein type I (NF1)]
700001	COLOBECTAL MUTANT CANCER PROTEIN

TABLE 6 (CONT)

[L25610] [M64349] [M90813]		O and Manual
[L25610] [M64349] [M90813] [K03214;		
[L25610] [M64349] [M90813] [K03214;		15
[L25610] [M64349] [M90813]	OC / I QIMI	MI ECOLO CELL NOCLEAN DIFFERENTIALIDIA ANTIGEN
[L25610] [M64349] [M90813] [K03214;	M81840	
[L25610] [M64349] [M90813] [K03214;	M83234	Y BOX BINDING PROTEIN-1 [Nuclease-sensitive element DNA-binding protein]
[L25610] [M64349] [M90813] [K03214;	U02082	GUANINE NUCLEOTIDE REGULATORY PROTEIN TIM1
[L25610] [M64349] [K03214;	U03056	HYALURONIDASE [tumor suppressor (LUCA-1)]
[L25610] [M64349] [M90813] [K03214;	U07236	PROTO-ONCOGENE TYROSINE-PROTEIN KINASE LCK [Lymphocyte-specific protein krosine kinase]
[M64349] [M90813] [K03214;	U09579; [L25610]	CYCLIN-DEPENDENT KINASE INHIBITOR 1 (MELANOMA DIFFERENTIATION ASSOCIATED PROTEIN 6) (MDA-6) (P21) (CDK-INTERACTING PROTEIN 1) (CIP1) (WAF1) (CDKN1A) (CDKN1) (SDI1) (PIC1) (CAP20)
SKI ONCOGENE SKI-NELATED ONCOGENE SNON WILMS TUMOR PROTEIN Cdc25A; M-PHASE INDUCER PHOSPY CYCLIN D3 PROTO-ONCOGENE (SERINE/INDOTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE PROTEIN SRC). C-REL PROTO-ONCOGENE PROTEIN SRC). C-REL PROTO-ONCOGENE PROTEIN SHOA SHB ADAPTOR PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HOCC25 [GUANINE NUCLEOTIDE	X07024	TRANSCRIPTION INITIATION FACTOR TFIID (250 KD SUBUNIT) [CG1 protein inv. in cell proliferation]
SKI-RELATED ONCOGENE SNON WILMS TUMOR PROTEIN cdc25A; M-PHASE INDUCER PHOSPI CYCLIN D3 PROHIBITIN RAF PROTO-ONCOGENE (SERINE/I PROTO-ONCOGENE TYROSINE-PRO [M90813] CYCLIN D2 [K03214; PROTO-ONCOGENE TYROSINE-PRO SRC). C-REL PROTO-ONCOGENE PROTEIN C-REL PROTO-ONCOGENE PROTEIN SHB ADAPTOR PROTEIN RHOA SHB ADAPTOR PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src H CDC25 [GUANINE NUCLEOTIDE	X15218	SKI ONCOGENE
WILMS TUMOR PROTEIN cdc25A; M-PHASE INDUCER PHOSPI CYCLIN D3 PROHIBITIN RAF PROTO-ONCOGENE (SERINE/I PROTO-ONCOGENE TYROSINE-PROTOL-1 (W64349) CYCLIN D1 (CYCLIN PRAD1) (BCL-1 (M90813) CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 [M90813] CYCLIN D2 SRC) C-REL PROTO-ONCOGENE PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HOCOCCES (GUANINE NUCLEOTIDE COCCES (GUANINE NUCLEOTIDE COCCES)	X15219	SKI-RELATED ONCOGENE SNON
CYCLIN D3 PROHIBITIN RAF PROTO-ONCOGENE (SERINET) PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE TYROSINE-PROTO-ONCOGENE PROTEIN SRC). C-REL PROTO-ONCOGENE PROTEIN RHOA SHB ADAPTOR PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HOCOCS [GUANINE NUCLEOTIDE] TIMOR SUPPRESSOR PROTEIN DC	X51630	WILMS TUMOR PROTEIN
CYCLIN D3 PROHIBITIN RAF PROTO-ONCOGENE (SERINE/TPROTO-ONCOGENE TYROSINE-PRC CYCLIN D1 (CYCLIN PRAD1) (BCL-1 CYCLIN D2 PROTO-ONCOGENE TYROSINE-PRC SRC). C-REL PROTO-ONCOGENE PROTEIL TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HC CDC25 [GUANINE NUCLEOTIDE	M81933	cdc25A; M-PHASE INDUCER PHOSPHATASE 1 (EC 3.1.3.48)
PROHIBITIN RAF PROTO-ONCOGENE (SERINE/I PROTO-ONCOGENE TYROSINE-PRC CYCLIN D1 (CYCLIN PRAD1) (BCL-1 CYCLIN D2 PROTO-ONCOGENE TYROSINE-PRC SRC). C-REL PROTO-ONCOGENE PROTEIN TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN (A Src H CDC25 (GUANINE NUCLEOTIDE TIMOR SUPPRESSOR PROTEIN DC	M92287	CYCLIN D3
RAF PROTO-ONCOGENE (SERINE/I PROTO-ONCOGENE TYROSINE-PRO CYCLIN D2 PROTO-ONCOGENE TYROSINE-PRO SRC). C-REL PROTO-ONCOGENE PROTEII TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN (A Src H CDC25 (GUANINE NUCLEOTIDE	S85655	PROHIBITIN
PROTO-ONCOGENE TYROSINE-PRC CYCLIN D1 (CYCLIN PRAD1) (BCL-1 CYCLIN D2 PROTO-ONCOGENE TYROSINE-PRC SRC). C-REL PROTO-ONCOGENE PROTEIL TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src H CDC25 [GUANINE NUCLEOTIDE	X03484	RAF PROTO-ONCOGENE (SERINE/THREONINE-PROTEIN KINASE)
CYCLIN D1 (CYCLIN PRAD1) (BCL-1 CYCLIN D2 PROTO-ONCOGENE TYROSINE-PRC SRC). C-REL PROTO-ONCOGENE PROTEII TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HC CDC25 [GUANINE NUCLEOTIDE	X16416	PROTO-ONCOGENE TYROSINE-PROTEIN KINASE ABL
CYCLIN D2 K03214; PROTO-ONCOGENE TYROSINE-PRO SRC). C-REL PROTO-ONCOGENE PROTEIL C-REL PROTO-ONCOGENE PROTEIL TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN A Src HC CDC25 GUANINE NUCLEOTIDE TUMOR SUPPRESSOR PROTEIN DC	X59798; [M64349]	CYCLIN D1 (CYCLIN PRAD1) (BCL-1 ONCOGENE)
K03214; PROTO-ONCOGENE TYROSINE-PRO SRC). C-REL PROTO-ONCOGENE PROTEIL TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN A Src H CDC25 (GUANINE NUCLEOTIDE TUMOR SUPPRESSOR PROTEIN DO TIMOR SUPPRESSOR PROTEIN DO		CYCLIN D2
C-RÉL PROTO-ONCOGENE PROTEIL TRANSFORMING PROTEIN RHOA SHB ADAPTOR PROTEIN (A Src H CDC25 (GUANINE NUCLEOTIDE TUMOR SUPPRESSOR PROTEIN DO	HT2291; [K03214; X03996]	PROTO-ONCOGENE TYROSINE-PROTEIN KINASE SRC (EC 2.7.1.112) (P60-SRC) (C-SRC).
SHB ADAPTOR PROTEIN RHOA SHB ADAPTOR PROTEIN [A Src HC CDC25 [GUANINE NUCLEOTIDE	X75042	C-REL PROTO-ONCOGENE PROTEIN
SHB ADAPTOR PROTEIN [A Src HC CDC25 [GUANINE NUCLEOTIDE		- 1
SHB ADAPT CDC25 TUMOR SUF	L.25080	I KANSFUHMING PHUTEIN HHUA [proto-oncogene rnoA, muttidrug resistance protein]
CDC25	X75342	APT
	L26584	
	X76132	TUMOR SUPPRESSOR PROTEIN DCC

TABLE 6 (CONT)

<u> </u>	127211	COLOR IN PERCENTANT VINASE A INCIDITOD A CONTANT (DAS INICA) (DAS INICA)
		CTCLIN-DEPENDEN! NIMASE 4 INTIDITION A (CDN+1) (TIC-1141/4) (TIC-1141/4)
		(MULTIPLE TUMOR SUPPRESSOR 1) (MTS1). (CDKN2A)
2	M13228	N-MYC PROTO-ONCOGENE PROTEIN
2	M15400	RETINOBLASTOMA-ASSOCIATED PROTEIN [retinoblastoma susceptibility]
2	M15990	PROTO-ONCOGENE TYROSINE-PROTEIN KINASE YES
2	M19720	L-MYC-2 PROTEIN
2	M19722	PROTO-ONCOGENE TYROSINE-PROTEIN KINASE FGR (EC 2.7.1.112) (P55-FGR) (C-
		FGR).
4	M73812	CYCLIN E (G1/S-SPECIFIC)
-	M74088	ADENOMATOUS POLYPOSIS COLI PROTEIN
-	U25994	TYROSINE-PROTEIN KINASE LYN [cell death protein RIP]
<u></u> 	U40343; [U20498]	CYCLIN-DEPENDENT KINASE 4 INHIBITOR D (P19-INK4D).
	U43746	BREAST CANCER TYPE 2 SUSCEPTIBILITY PROTEIN
	X02751	TRANSFORMING PROTEIN P21 [N-ras]
	X16706	FRA-2 [fos-related antigen 2]
	X16707	FRA-1 [fos-related antigen 1]
	X51521	EZRIN [Villin 2]
	X56681	TRANSCRIPTION FACTOR JUN-D
	X59932	TYROSINE-PROTEIN KINASE CSK [C-SRC-kinase]
	X86779	FAST KINASE
	X87838	BETA-CATENIN
	Z29090	PHOSPHATIDYLINOSITOL 3-KINASE CATALYTIC SUBUNIT ALPHA ISOFORM
	M14745	BCL2
	D38305	
	L16464	ETS-RELATED PROTEIN PE-1 [ETS oncogene (PEP1)]
	L29216	PROTEIN KINASE CLK (CLK2)
	L29220	PROTEIN KINASE CLK (CLK3)
	L29222	PROTEIN KINASE CLK (CLK1)
	U10564	CDK TYROSINE 15-KINASE WEE1Hu

TABLE 6 (CONT)

9	GenBank #	Gene Name
<u> </u>	U22398	CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE INHIBITOR 1
		r3/) (r3/nir2)
2_	U24166	EB1
<u>n</u>	U26710	PROTO-ONCOGENE C-CBL
<u> </u>	U33841	ATAXIA TELANGIECTASIA (ATM)
)	U35735	RACH1
0	U40282	INTEGRIN-LINKED KINASE (ILK) [MIXED LINEAGE KINASE 2]
3	U41816	C·1
7	U43408	FOCAL ADHESION KINASE [tyrosine kinase (Tnk1)]
٦.	U57456	MOTHERS AGAINST DPP PROTEIN (chromosome 4 Mad homolog Smad1; transforming
		growin ractor-pera signaling protein-1 (usp-1)
<u></u>	00809	- 1
	U61262	TUMOR SUPPRESSOR PROTEIN DCC [neogen.n]
	U63139	DNA REPAIR PROTEIN RAD50
2	M81934; [S78187]	
	U17075; [L36844]	CYCLIN-DEPENDENT KINASE 4 INHIBITOR B (P14-INK4B) (P15-INK4B) (MULTIPLE
		I UMOH SUPPHESSON Z) (MISZ) (CONNZE).
	U84119	LACTOFERRIN (DELTA)
	X74262	RBA/p48
	X85133	RBQ1 retinoplastoma binding protein
	Z29083	5T4 ONCOFETAL ANTIGEN
	123959	E2F-related transcription factor (DP-1)
	125676	SERINE/THREONINE PROTEIN KINASE PITALRE
	L26081	semaphorin III
	L37882	frizzled
	L20861	Wnt-5a
	M29039	jun B TRANSACTIVATOR
	M34065	cdc25C; M-PHASE INDUCER PHOSPHATASE 3 (EC 3.1.3.48).
	M73980	Notch1
	M95712	ral,b-
	M99437	notch group protein (N)
	U15642	E2F-5
	U33920	semaphorin V

TABLE 6 (CONT)

:	1149318	
		(rizzled 5
	1,0101	dishavalled homolog (DVI)
: : : :	Т.	United the second of the secon
	U49262; [U75651]	dishevelled (DVL) + dishevelled 3 (DVL3)
		MATE OF IDE
	X07876	WNIXONIN
	L40027	glycogen synthase kinase 3
	X66360	SERINE/THREONINE-PROTEIN KINASE PCTAIRE-2
	X66362	SERINE/THREONINE PROTEIN KINASE PCTAIRE-3
	X66363	SERINE/THREONINE-PROTEIN KINASE PCTAIRE-1
	X74594	RB2/p130
	X85134	RBQ-3
	Z71621	Wnt-13
	AB000220	semaphorin E
	AF001954	growth inhibitor p33ING1 (ING1)
	AF007111	MDM2-like p53-binding protein (MDMX)
	D89667	C-myc binding protein
	U29343	HYALURONAN RECEPTOR (RHAMM)
	U66469	p53-dependent cell growth regulator CGR19
	U76638	BRCA1-ASSOCIATED RING DOMAIN PROTEIN
	U82169	frizzled homolog (FZD3)
	U84401	smoothened
	U90875	cytotoxic ligand TRAIL receptor
	U95299	Notch4
	Y11416	p73, a monoallelically expressed p53-related protein
	X91940	WNT-8B
	X97057	WNT-10B
	Y10479	E2F-3
	Y11306	beta catenin/TCF-4
	U38276	SEMAPHORIN-1
	U77493	Notch2
	K00650	C-fos
	X53795	CD82 ANTIGEN (INDUCIBLE MEMBRANE PROTEIN R2) (C33 ANTIGEN) (IA4) [METASTASIS SUPPRESSOR KANGAI 1) (SUPPRESSOR OF TUMORIGENICITY-6).
	L38518	sonic hedgehog (SHH)
:	M54968	K-BAS, ONCOGENE

TABLE 6 (CONT)

GenBank #	Gene Name
M63167	Akt1 (rac protein kinase alpha, protein kinase B, c-Akt)
S57153; S57160	RBP1(RETINOBLASTOMA-BINDING PROTEIN)
U23435; U31089	Abi interactor 2 (Abi-2) + Abl binding protein 3 (AbiBP3) [ArgBPIB]
M96577	E2F-1 pRB-binding protein
U24163; [U91903;	U24163; [U91903; frizzled-related FrzB (Fritz) (frezzled (fre))
L05148	TYROSINE-PROTEIN KINASE ZAP-70 (EC 2.7.1.112) (70 KD ZETA-ASSOCIATED PROTEIN) (7AP70)
M97935	SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 1-ALPHA/BETA
U10087 X58957	TYROSINE-PROTEIN KINASE BTK (EC 2.7.1.112) (BRUTON'S TYROSINE KINASE)(AGAMMAGLOBULINAEMIA TYROSINE KINASE) (ATK) (B CELL PROGENITOR KINASE) (BPK) (BTK) (AGMX1)
AF016268	death receptor 5 (DR5)
M35296	TYROSINE-PROTEIN KINASE ABL2 (EC 2.7.1.112) (TYROSINE KINASE ARG) (ABLL)
U18671 M97934	SIGNAL TRANSDUCER AND ACTIVATOR OF TRANSCRIPTION 2 (P113) (STAT2)
U47686	SIGNAL TRANSDUCER AND TRANSCRIPTION ACTIVATOR 58 (STAT58)
M80629	CDC2-RELATED PROTEIN KINASE CHED
S66431	RBP2 retinoblastoma binding protein
U04045; [L47583]	DNA MISMATCH REPAIR PROTEIN MSH2
U29656	DR-NM23
U43148	patched homolog (PTC)
J02958	MET
U49089	neuroendocrine-dig (NE-dig) a novel human homolog of the Drosophila discs large (dig) fumor suppressor protein interacting with the APC protein
U54777	DNA MISMATCH REPAIR PROTEIN MSH6 (mutS - ALPHA 160 KD SUBUNIT) (G/T MISMATCH BINDING PROTEIN) (GTBP) (GTMBP) (P160)
X66358	SERINE/THREONINE-PROTEIN KINASE KKIALRE

Cell-Cell Interaction Array

5

In the cell-cell interaction array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that are associated with cell-cell interaction, e.g. cell-cell signaling. In a specific cell-cell interaction array of interest, the spots are as provided in Table 7.

TABLE 7

Cena		l
M32315		ᆱ
X01394		CTOR (TNFa)
D12614		LYMPHOTOXIN-ALPHA [formerly tumor necrosis factor beta (TNF-beta)]
M12807		<u>آ</u> ر
M14648		VITRONECTIN RECEPTOR ALPHA [Integrin, alpha V; antigen CD51]
X75208		밁
X74764		TYROSINE-PROTEIN KINASE CAK [Tyrosine kinase, receptor TKT]
M18391	161	TYROSINE-PROTEIN KINASE RECEPTOR EPH
0088	39 [M83246;	<u>U08839 M83246; UROKINASE PLASMINOGEN ACTIVATOR SURFACE RECEPTOR, GPI-ANCHORED</u>
X51675	751	FORM PRECURSOR (U-PAR) (MONOCYTE ACTIVATION ANTIGEN MO3) (CD87
		ANTIGEN)
M33294		TUMOR NECROSIS FACTOR RECEPTOR [Tumor necrosis factor receptor 1 (55kD)]
Y00285		CATION-INDEPENDENT MANNOSE-6-PHOSPHATE RECEPTOR Insuline-like grown
		factor receptor II, IGFR-2]
L07414		CD40
L08096;		CD27 (CD70 ANTIGEN)
[866333]	339]	
L09753	53	CD30
M35410	410	=
M63928	928	اہ
M67454	454	닉
M83554	554	
X60592	192	CD40L RECEPTOR [Cdw40 nerve growth factor receptor-related B-lymphocyte activation
		molecule
D138	D13866 [D14705	ALPHA-CATENIN (CADHERIN-ASSOCIATED PROTEIN) (ALPHA E-CATENIN)
1238	123805; 122080]	
D25303;	303;	integrin alpha9
[124	L24158]	
J03132	132	INTERCELLULAR ADHESION MOLECULE-1
J04536	336	LEUKOSIALIN (sialophorin (CD43))
L113	L11353; Z22664;	MERLIN (SCHWANNOMIN) (moesin-ezrin-radixin-like protein)(neurofibromatosis 2)
X726	X72657; L27133	
L13616	516	Focal adhesion kinase
L14837	837	TIGHT JUNCTION PROTEIN ZO-1
L16785;	785;	NUCLEOSIDE DIPHOSPHATE KINASE B (EC 2.7.4.6) (NDK B) (NDP KINASE B) (NM23-H2)
LEAD	114250811	(C.MYC PIIRINE-BINDING TRANSCRIPTION FACTOR PUF).

TABLE 7 (CONT)

1 20815	
1.25259	T LYMPHOCYTE ACTIVATION ANTIGEN CD86 [CD28 antigen ligand 2, B7-2 antigen]
L34774	opioid binding cell adhesion molecule
M15476	UROKINASE-TYPE PLASMINOGEN ACTIVATOR PRECURSOR (EC 3.4.21.7.3) (UPA) (UP
M15518; [TISSUE-TYPE PLASMINOGEN ACTIVATOR PRECURSOR (EC 3.4.21.68) (T-PA) (T-PA)
X07393; M1	X07393; M18182] PLASMINOGEN ACTIVATOR).
M18082:[PLASMINOGEN ACTIVATOR INHIBITOR-2, PLACENTAL (PAI-2) (MONOCYTE ARG-
102685	
M21097	<u></u>
M23197	CD33 MYELOID CELL SURFACE ANTIGEN Unreferritation anugen (CD33)
M28882	m i
M30257	崱
M30640	E-SELECTIN (Endothelial leucocyte adhesion molecule i (ELAM I)
M34064 [X57548;	57548; CADHERIN-2 (N-CADHERIN)
X54315; S42303]	[2303]
	CO22 B.CELL DIFFERENTIATION ANTIGEN
MORPO	CD44 ANTIGEN HEMATOPOIETIC FORM [Cell adhesion molecule (CD44)]
M63648	billous pemphigoid antigen
M74387	LICAM
M74777	1
U01160	
U03056	ASE [turnor su
U07819	CONTACTIN (Contactin 1 (CNTN1))
U15979	DELTA-LIKE PROTEIN [dlk]
X16841	N-CAM (NEURAL CELL ADHESION MOLECULE, PHOSPHATIDY LINOSITOL-LINNED
	ISOFORM; CD56]
X70326	Ì
X74979	TYROSINE-PROTEIN KINASE CAK [EDDR1; TRK E]
Z26317 [S64273]	desmoglein 2
172080	TRANSFORMING PROTEIN RHOA [proto-oncogene rhoA, multidrug resistance protein]
X76132	200
102703	TO A TELET MEMBRANE GLYCOPROTEIN IIIA

TABLE 7 (CONT)

GenBank #	(CELL INTERACTION (Gene Names)
304145	INTEGRIN ALPHA M [Neutrophil adherence receptor alpha-M subunit; Complement
	component receptor 3, alpha; also known as CD11b (p170), macrophage antigen alpha
	polypeptide
105633	integrin beta5
L12002;	integrin alpha4
[X16983]	
125851	integrin alphaE
L36531	integrin alpha8
M15395	LEUKOCYTE ADHESION PROTEIN (CELL SURFACE ADHESION GLYCOPROTEINS LFA-
	1, CR3 AND P150,95, BETA-SUBUNIT]
M28249:	integrin alpha2 [very late antigen-2 (vla-2)/collagen receptor alpha-2 subunit]
[X17033]	A HIGGI GII HIGGIGANA
M34480	INTEGRIN ALPHA 2B (PLATELET MEMBRANE GLYCOPHOLEIN IIB (GPIID); anugen
	CD41B
M35198	integrin beta6
M59911	integrin alpha3
M62880	integrin beta7
M73780	Integrin beta8
M81695	INTEGRIN ALPHA X (LEUKOCYTE ADHESION GLYCOPROTEIN P150,95 ALPHA CHAIN;
	antigen CD11C (p150))
X06256	딀
87670X	FIBRONECTIN RECEPTOR (BETA SUBUNIT) (INTEGRIN BETA 1)
X53586;	integrin alpha6
[X59512]	
X53587;	integrin beta4
[X52186]	
X68742	integrin alpha
X74295	integrin alpha7B
V00796	INTEGRIN ALPHA L (LEUKOCYTE ADHESION GLYCOPROTEIN LFA-1 ALPHA CHAIN;
	antigen CD11A (p180)]
D38122	FAS ANTIGEN LIGAND
M74088;	APC (DP2.5)
[M73548]	
U43522;	Protein tyrosine kinase Pyk2 (Cell adhesion kinase-beta, CAK-beta) (FAK2)
[L49207]	
X51521	Ezrin (cytovillin 2)

TABLE 7 (CONT)

10051	GenBank #	CELL INTERACTION (Gene Names)
[D10051]	X87838 [Z19054]	BETA-CATENIN
[J05471]	L11015	-BETA
;; ;; S5] S5] S6] S6] S7, [J05477]	U57059	FAS ANTIGEN LIGAND (TNF-related apoptosis inducing ligand TRAIL; Apo-2 ligand)
55 51 51 7, [J05471] 1, [D10051]	D45132	ANNEXIN I (zinc finger protein RIZ)
	M68516;	PLASMA SERINE PROTEASE INHIBITOR PRECURSOR (PCI) (PROTEIN C INHIBITOR)
	[102639]	(PLASMINOGEN ACTIVATOR INHIBITOR-3) (PAI3).
	U40282	ļ
	U43408	
	U60800	
	U61262	
	L11370	protocadherin 42
, [J05471] , [J105471]	X78817	RHO-GAP HEMATOPOIETIC PROTEIN C1 (P115) (KIAA0131).
[5] [5] [5] [5] [5] [5] [6] [6] [6] [7] [6] [7] [7] [7] [7] [7] [7] [7] [7] [7] [7	X85978	TAX1, AXONIN-1/TAQ1
[5] [105471] [D10051];	L11373	protocadherin 43
[5] [105471] [D10051];	X89576	MMP-17 (MT4-MMP)
[] [D1005471] [D10051];	Y00815	LAR
[] [D1005471] [D10051];	230183	TIMP-3 (mitogen-inducible gene 5, mig-5)
[J05471] [D10051]	Z35227	ras-like small GTPase TTF
[J05471] [D10051]	D26512,	MMP-14 (MT1-MMP)
[D10051]	[X83535]	
[D10051]	D31784	CADHERIN-6
[D10051]	D50477	MMP-16 (MT3-MMP)
J03210, [J05471] MMP-2 (gelatinase A) J05070, [D10051] MMP-9 (gelatinase B) J05556 MMP-8 (collagenase-2) L2068B rho GDP-dissociation inhibitor protein 2 (Ly-GDI) L26081 semaphorin III L34056 CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN) L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN, NEURAL TYPE	D83542	CADHERIN-14 MUSCLE-CADHERIN PRECURSOR (M-CADHERIN) (CADHERIN-14) ICADHERIN-15
J055070, [D10051] MMP-9 (gelatinase B) J05556 MMP-8 (collagenase-2) L2068B the GDP-dissociation inhibitor protein 2 (Ly-GDI) L26081 semaphorin III L34056 CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN) L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN) N-CADHERIN, NEURAL TYPE	J03210, [J05471] MMP-2 (gelatinase A)
J05556 MMP-8 (collagenase-2) L20688 tho GDP-dissociation inhibitor protein 2 (Ly-GDI) L26081 semaphorin III L34056 CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN) L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL TYPE	J05070, [D1005	1] MMP-9 (gelatinase B)
L26688 the GDP-dissociation inhibitor protein 2 (Ly-GDI) L26081 semaphorin III L26081 CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN) L34056 L34056 L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL TYPE	305556	MMP-8 (collagenase-2)
126081 semaphorin III 134056 CADHERIN CADHERIN	120688	rho GDP-dissociation inhibitor protein 2 (Ly-GDI)
L34056 CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN) L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL TYPE	L26081	semaphorin III
L34057; [L33477] CADHERIN-12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL 1YPE	L34056	CADHERIN-11 (OSTEOBLAST-CADHERIN) (OB-CADHERIN)
	L34057; [L3347	7]CADHERIN-12 (BR-CADHERIN) (N-CADHERIN 2) (CADHERIN, NEURAL TYPE, 2)

TABLE 7 (CONT)

_	GenBank #	CELL INTERACTION (Gene Names)
	L34058;	CADHERIN-13 T-CADHERIN PRECURSOR (TRUNCATED-CADHERIN) (H-CADHERIN)
	(US9289;	(HEART-CADHERIN)
	U59288 L34059	CADHERIN-4 RETINAL-CADHERIN PRECURSOR (R-CADHERIN) (R-CAD)
:	L34060	CADHERIN-8
	M23410	PLAKOGLOBIN (DESMOPLAKIN III)
	M94151	ALPHA-CATENIN RELATED PROTEIN (CATENIN ALPHA-2)
	U24152	SERINE/THREONINE-PROTEIN KINASE PAK-ALPHA (EC 2.7.1) (P65-PAK) (P21-
		ACTIVATED KINASE) (ALPHA-PAK)
	U24153	p21-activated protein kinase (Pak2)
	U33920	semaphorin V
	U43318	frizzled 5
	X04429	PLASMINOGEN ACTIVATOR INHIBITOR-1 PRECURSOR, ENDOTHELIAL (PAI-1)
	X13916	LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 1 PHECURSOR (LRP) (ALPHA-2-MACROGLOBULIN RECEPTOR) (A2MR)
	X14787	THROMBOSPONDIN 1 PRECURSOR
	L40027	glycogen synthase kinase 3
	X54412	collagen type IX alpha-1
	X56654	desmoglein type 1
	X56807	DSC2 mRNA for desmocollins type 2a and 2b
	X61587	rhoG
	X63629	CADHERIN-3 PLACENTAL-CADHERIN PRECURSOR (P-CADHERIN)
	X69550	rho GDP-dissociation Inhibitor 1
	X75308	MMP-13 (collagenase-3)
	X78565	TENASCIN-C
	X79981;	CADHERIN-5 VASCULAR ENDOTHELIAL-CADHERIN PRECURSOR (VE-CADHERIN) (784
	[36765X]	ANTIGEN) (CD144 ANTIGEN).
	M11313	ALPHA-2-MACROGLOBULIN PRECURSOR (ALPHA-2-M)
	X95282	Rho8 protein
	X95456	Rho7 protein
	Y07923	Rho6 protein
	Z13009	CADHERIN-1(E-CADHERIN) (UVOMORULIN) (CAM 120/80)
	Z15009	laminin
	Z48482	MMP-15 (MT2-MMP)
	AB000220	semaphorin E
!	0000	

TABLE 7 (CONT)

Genbank # Genbank # Genbank # D85815 D85815 D85815 D29343 D72661 U76456 U76456 U76456 U76456 U38276 U943576 U943576 U0943570 U02570 X05934 X05954 X05954 X05954 X05934 X05954	rhoHPTI AMP-19 ININJURIN-1 TIMP-4 GDI-dissociation inhibitor RhoGDIgamma MMP-19 INTS3-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 Iunatic fringe CDCA2 GTPase-activating protein
M24795 M29343 M29343 M29343 M2795 M2610 M2621 M26221 M26221 M11306 M13264 M13364 M38276 M	
AF000974 U29343 M24795 M24795 U72661 U76456 U82532 X92521 Y07604 Y11306 U38276 U94354 U02570	
W29343 W24795 W24795 U7661 U76456 U82532 X92521 Y07604 Y11306 U38276 U94354 U02570	
M24795 U72661 U76456 U82532 X92521 Y07604 Y11306 U38276 U94354 U02570 X05331	PLATELET GLYCOPHOTEIN IV (GTIN) (GTIN) (GTIN) (CDS) CANTELY (COPHOTEIN) NINJURIN-1 TIMP-4 GDI-dissociation inhibitor RhoGDIgammma MMP-19 nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 Iunatic fringe
U72661 U76456 U82532 X92521 Y07604 Y11306 U38276 U94354 U02570 X062319	NINJURIN-1 TIMP-4 GDI-dissociation inhibitor RhoGDIgammma GDI-dissociation inhibitor RhoGDIgammma MMP-19 nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 Iunatic fringe CDCA2 GTPase-activating protein
V76456 U82532 X92521 Y07604 Y11306 U38276 U94354 U02570 X05331	TIMP-4 GDI-dissociation inhibitor RhoGDIgammma MMP-19 nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 lunatic fringe CDC42 GTPase-activating protein
V02532 X92521 Y07604 Y11306 U38276 U94354 U02570 X05331	GDI-dissociation inhibitor RhoGDIgammma MMP-19 nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 lunatic fringe CDC42 GTPase-activating protein
X92521 Y07604 Y11306 U38276 U94354 U02570 X05331	MMP-19 nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC 2.7.4.6) (NUCLEOSIDE 5'- DIPHOSPHATE PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 lunatic fringe CDC42 GTPase-activating protein
Y07604 Y11306 U38276 U94354 U02570 X05331	nm23-H4; NUCLEOSIDE-DIPHOSPHATE KINASE (EC.2.7.4.0) (NOCLEOSIDE OF PHOSPHOTRANSFERASE) (NDK). beta catenin/TCF-4 SEMAPHORIN-1 lunatic fringe CDC42 GTPase-activating protein
V11306 U38276 U94354 U02570 X05331	beta catenin/TCF-4 SEMAPHORIN-1 Iunalic fringe CDC49 GTPase-activating protein
V11306 U38276 U94354 U02570 X05199	SEMAPHORIN-1 Iunatic fringe
U94354 U94354 U02570 X05199	Iunatic fringe
U02570 X05199	Concas activating protein
002570 X05199	COCA CI Pase-activating Diologic
X05199	12 CO CO CO CO CO CO CO CO CO CO CO CO CO
V0E231	PLASMINOGEN PHECURSON (EC 3.4.51.1)
7300	MMP-1 (collagenase-1)
X53795	CD82 ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN (INDUCIBLE MEMBRANE PROTEIN RZ) (USS ANTIGEN RZ)
	(METASTASIS SUPPHESSON KANGALI) (SOFT INCOMES
L38517	indian hedgehog protein (HHI)
M31470	
M34189	
X83929;	desmocollin type 3 + desmocollin type 4
[D17427	
123808	MMP-12 (metalloelastase)
125081	IrhoC (H9); SMALL GTPase (moC)
M29870;	
[M31467]	
M64595;	RAS-RELATED C3 BOTULINUM TOXIN SUBSITIALE 1 (PZI-HACZ)
[M29871]	
X05232	MMP-3 (stromelysin-1)
X06820	
X07820,	MMP-10 (stromelysin-2)
[M30461]	
X72925	desmocollin type 1

TABLE 7 (CONT)

GenBank#	CELL INTERACTION (Gene Names)
X94991;	Zyxin + Zyxin-2
[X95735]	
U52111	PLEXIN
M38690	CD9
M54995; M38	M54995; M38441 PLATELET BASIC PROTEIN PRECURSOR (PBP) (CONTAINS: CONNECTIVE-TISSUE
	BETA-THROMBOGLOBULIN (BETA-TG), NEUTROPHIL-ACTIVATING PEPTIDE 2 (NAP-2))
1 20471	extracellular matrix metalloproteinase inducer EMMPRIN
M57730 M37476	
	(LERK-1) (IMMEDIATE EARLY RESPONSE PROTEIN B61) (TUMOR NECROSIS FACTOR)
	ALPHA-INDUCED PROTEIN 4).
0.007695	KINASE RECEPTOR HTK).
U09304	EPHRIN-B1 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2)
	(LERK-2) (ELK LIGAND PRECURSOR) (ELK-L).
U41766	metalloprotease/disintegrin/cysteine-rich protein precursor (MDC9)
U26403	EPHRIN-A5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7)
	(LERK-7) (AL-1).
AF035752	caveolin-2
U32114	to UNIVERSITY DE LA COMPANIE DE LA C
U66406	EPHRIN-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8)
	(LERK-8) (EPH-HELAI ED RECEP IOR I HANSMEMBRAINE LIGAND ELN-ES)
X95425	EPHRIN TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN
-1-	KINASE RECEPTOR ERK-1) (EPH HOMOLOGI NIKASE-1) (NECEPTOR TROTEIN)
Z18951 S49856	1
L38734	Π
•	(LERK-5) (HTK LIGAND) (HTK-L).
L40636	EPHRIN TYPE-B RECEPTOR 1 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN
	KINASE RECEPTOR EPH-2) (NET).
L41939	EPHRIN TYPE-B RECEPTOR 2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN EPH-
M16591	TYROSINE-PROTEIN KINASE HCK (EC 2.7.1.112) (P59-HCK AND P60-HCK)
	(HEMOPOIETIC CELL RINASE).

TABLE 7 (CONT)

GenBank #	CELL INTERACTION (Gene Names)
M59371 M36395	EPHRIN TYPE-A RECEPTOR 2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN
MEZOEO	AI PHA 2-MACROGI OBIJI IN RECEPTOR-ASSOCIATED PROTEIN PRECURSOR (ALPHA-
	2-MRAP) (LOW DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN- ASSOCIATED
	PROTEIN 1) (RAP)
M77830	desmoplakin l
M86826	IGF BINDING PROTEIN ACID-LABILE SUBUNIT
M99487	PROSTATE-SPECIFIC MEMBRANE ANTIGEN (PSM)
U04441	LOW-DENSITY LIPOPROTEIN RECEPTOR-RELATED PROTEIN 2 (MEGALIN)
	(GLYCOPROTEIN 330) (FRAGMENT)
U11690	PUTATIVE RHO/PAC GUANINE NUCLEOTIDE EXCHANGE FACTOR(RHO/PAC GEF)
	(FACIOGENITAL DYSPLASIA PROTEIN)
U14588	Paxillin
U16296	T-lymphoma invasion and metastasis inducing TIAM1
U29656	DR-NM23
U32907	P37NB
U35113	METASTASIS-ASSOCIATED MTA1
U37139	beta 3-endonexin
U43195	Rho-associated, coiled-coil containing protein kinase p160ROCK
U43527	malignant melanoma metastasis-suppressor (KiSS-1) gene
U49089	neuroendocrine-dig (NE-dig) a novel human homolog of the Drosophila discs large (dig) tumor
	suppressor protein interacting with the APC protein
U53786	envoplakin (EVPL)
U59752	cytohesin-1; Sec7p-like protein
X03124	TIMP-1 (erythroid potentiating activity, EPA)
X07819	MMP-7 (matrilysin)
X17620	NUCLEOSIDE DIPHOSPHATE KINASE A (EC 2.7.4.6) (NDK A) (NDP KINASE A) (TUMOR
	METASTATIC PROCESS-ASSOCIATED PROTEIN) (METASTASIS INHIBITION FACTOR
	NM23) (NM23-H1).
105593	TIMP-2 (MI)
X57766	MMP-11 (stromelysin-3)

Cytokine and Cytokine Receptor Array

5

In the cytokine and cytokine receptor array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that express cytokines or cytokine receptors. In a specific cytokine and cytokine receptor array of interest, the spots are as provided in Table 8.

TABLE 8

Cenbank	
20070	
X01992	IN EDITOR CANAMIC
J04156	
X01057	INTERLEUKIN-2 RECEPTOR ALPHA CHAIN
A14844	INTERLEUKIN-2
M29366	PROTEIN-TYROSINE KINASE RECEPTOR ERBB-3 [Epidermal growth factor receptor (awan
	erythroblastic leukemia viral (v-erb-b) oncogene homolog)]
X04434	INSULIN-LIKE GROWTH FACTOR I RECEPTOR
M29645	medin A]
X03663	MACROPHAGE COLONY STIMULATING FACTOR I RECEPTOR [c-fms proto-oncogene]
	THE SECRET SECTION REPORTED A PRECINENCE MINOR NECROSIS FACTOR
M32315;	BINDING PROTEIN 2) (TBPII) (P80) (TNF-R2) (P75) (CD1208) (TNFR2) (TNFBR).
X02811:	PLATELET-DERIVED GROWTH FACTOR, B CHAIN PRECURSOR (PDGF B-CHAIN)
[X02744	(PDGF-2) (BACAPLERMIN) (C-SIS)
M12783	
X02851	INTERLEUKIN-1 ALPHA
K02770	INTERLEUKIN IL-1BETA
M14743;	INTERLEUKIN-3 PRECURSOR (IL-3) (MULTIPOTENTIAL COLONY-SIMULA INC.
[M17115]	FACTOR) (HEMATOPOIETIC GROWTH FACTOR) (P-CELL STIMULATING FACTOR)
	(MAST-CELL GROWTH FACTOR) (MCGF) (IL3).
M13982	INTERLEUKIN-4
X04602;	
[M14584]	(INTERFERON BETA-2) (HYBRI
X01394	-
D12614	LYMPHOTOXIN-ALPHA (formerly tumor necrosis factor beta (INF-beta)
M20566	INTERLEUKIN-6 RECEPTOR ALPHA CHAIN
X04688;	INTERLEUKIN IL-5 (8 CELL DIFFERENTIATION FACTOR I) (T-CELL REPLACING
[103478]	FACTOR) (EOSINOPHIL DIFFERENTIATION FACTOR)
M28622	
M11220	GRANULOCYTE-MACROPHAGE COLONY-STIMULATING FACTOR [GM-CSF]
K03222	TRANSFORMING GROWTH FACTOR ALPHA
100000	LEUKOCYTE INTERFERON ALPHA
[100207	
X02812	1
X03438	GRANULOCYTE COLONY-STIMULATING FACTOR [G-CSF]
M19154	۷
X04571	H FACTOR KIDNEY [EGF]
J03171	HUIEN- ALPHA-REC (INTERFERON ALPHA-BETA RECEPTOR ALPHA CHAIN)
M57627	
0303011	NIATO CHOCOTO CHOM SETA CHAIN

TABLE 8 (CONT)

	M74782	INTERLEUKIN-3 RECEPTOR ALPHA CHAIN
!	X52425	NITERI EUKIN-4 RECEPTOR ALPHA CHAIN
	1475914	INTER EUKIN-S RECEPTOR ALPHA CHAIN
	X77722	INTERFEBON ALPHA-BETA RECEPTOR BETA CHAIN
	X72755	GAMMA INTERFERON INDUCED MONOKINE (Humig)
	D11086	CYTOKINE RECEPTOR COMMON GAMMA CHAIN [Interleukin 2 receptor gamma chain]
	M20132	ANDROGEN RECEPTOR
	M73238	PACTOR
	303143	INTERFERON GAMMA RECEPTOR ALPHA CHAIN
	M60459	ERYTHROPROTEIN RECEPTOR
	L00587	1
	M62424	THROMBIN RECEPTOR [Coagulation factor II (thrombin) receptor]
	L07594	TRANSFORMING GROWTH FACTOR-BETA TYPE III RECEPTOR
	M84747	INTERLEUKIN-9 RECEPTION
	U00672	INTERLEUKIN-10 RECEPTOR
	M14764	LOW-AFFINITY NERVE GROWTH FACION HECEPION
	X60957	TYROSINE-PROTEIN KINASE RECEPTOR TIE-1 PRECORSOR (EC 2.7.1.1.12).
	100000	VANCET AS ENDOTHER IA GROWTH FACTOR RECEPTOR 3 PRECURSOR (EC
	X682U3; [X69878:	2.7.1.112) (VEGFR-3) (TYROSINE-PROTEIN KINASE RECEPTOR FLT4, CLASS III).
	U43143	
	M16552	THROMBOMODULIN
	M87290	ANGIOTENSIN II RECEPTOR TYPE-1A
		TYROSINE-PROTEIN KINASE RECEPTOR ETK1
	M76673	FMLP-RELATED RECEPTOR I
	M97675	TRANSMEMBRANE RECEPTOR ROR1
!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!!	L04947;	VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR 2 PRECURSOR (EC
	[X61656]	2.7.1.112) (VEGFR-2) (KDR) (KINASE INSER! DOMAIN HECELIUM) (FINALMENT)
	M91196	INTERFERON CONSENSUS SEQUENCE BINDING PROTEIN DINA-BINGING PINCEIN
	X75208	TYROSINE-PROTEIN KINASE RECEPTOR EPH-3
	U05012	- 1
	X74764	TYROSINE-PROTEIN KINASE CAK [Tyrosine kinase, receptor TK1]
	K03193;	EPIDEAMAL GROWTH FACTOR RECEPTOR PRECURSOR (EC 2.7.1.112). (EGFH)
	(X00588;	(ERBB1)
	X00663;	
	U48722]	DI ATELET ACTIVATING FACTOR RECEPTOR
	M18391	TYROSINE-PROTEIN KINASE RECEPTOR EPH
:	A09781	INTERFERON-GAMMA RECEPTOR
	1112140	TYROSINE KINASE RECEPTOR TRK-B

TABLE 8 (CONT)

	M86492	Gene rame
	M86492	
-		-1
	L07868	ERBB4 (EPIDERMAL GROWTH FACTOR RECEPTOR)
	M27492	INTERLEUKIN-1 RECEPTOR TYPE I
	M33294	ı
==	M37435	MACROPHAGE COLONY STIMULATING FACTOR-1 [M-CSF]
	M11730	ERBB-2 RECEPTOR PROTEIN-TYROSINE KINASE
	D10923	HM74 PROBABLE G PROTEIN-COUPLED RECEPTOR HM74]
	D10924	HM89 [PROBABLE G PROTEIN-COUPLED RECEPTOR LCR1 HOMOLOG]
	D10925	HM145 [C-C CHEMOKINE RECEPTOR TYPE 1]
	D14012	HEPATOCYTE GROWTH FACTOR ACTIVATOR
	D16431	HEPTOMA-DERIVED GROWTH FACTOR
	D30751;	BONE MORPHOGENETIC PROTEIN 4 (BMP-2B)
	[M22490]	
-	J03358	낕
	J04130	MACROPHAGE INFLAMMATORY PROTEIN 1-BETA [Activation (Act-2)]
	J05081	ENDOTHELIN-3
	L06139	TYROSINE-PROTEIN KINASE RECEPTOR TIE-2 PRECURSOR (EC 2.7.1.112) (TYROSINE
		PROTEIN KINASE RECEPTOR TEK) (P140 TEK) (TUNICA INTERNA ENDOTHELIAL CELL
	L06622	ļ
	L06623	ENDOTHELIN B RECEPTOR (EDNHB)
	L06801	INTERLEUKIN-13
	L07414	
	L08096	
	L08187	CILIARY NEUROTROPHIC FACTOR RECEPTOR ALPHA (cytokine receptor EB13)
	109753	CD30
	L,12260;	RECOMBINANT GLIAL GROWTH FACTOR + NEU DIFFERENTIATION FACTOR +
	.U02326;	HEREGULIN
	M94165	- (
	L12261	HEREGULIN ALPHA [Recombinant glial growth factor]
	L15344	INTERLEUKIN IL-14
	L36052;	THROMBOPOIETIN PRECURSOR (MEGAKARYOCYTE COLONY STIMULATING
	[L36051;	FACTOR) (C-MPL LIGAND) (ML) (MEGAKARYOCYTE GROWTH AND DEVELOPMENT
	U11025	FACTOR) (MGDF) (THPO)
	M10051	INSULIN RECEPTOR
	M21121	RANTES PROTEIN T-CELL SPECIFIC
	M21574	PLATELET-DERIVED GROWTH FACTOR RECEPTOR ALPHA
	M21616	PLATELET-DERIVED GROWTH FACTOR RECEPTOR BETA
	M22488; [U50330]	BONE MORPHOGENETIC PROTEIN 1 (procollagen C-proteinase) (pCP-2)
	M22489	BONE MORPHOGENETIC PROTEIN 2A

TABLE 8 (CONT)

9	1 · · · · · · · · · · · · · · · · · · ·	
5	Genbank #	CENTE NAME
Σ	M22491	1
Ž	M23452	MACROPHAGE INFLAMMATORY PROTEIN 1-ALPHA [GOS19-1]
W.	M24545	MONOCYTE CHEMOTACTIC PROTEIN 1
<u>X</u>	M25667	NEUROMODULIN (Neuronal growth protein 43 (GAP-43))
M.	M27288	ONCOSTATIN M
×	M30704	AMPHIREGULIN [schwannoma-derived growth factor]
×	M31145	INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 1
×	M31165	TUMOR NECROSIS FACTOR-INDUCIBLE PROTEIN TSG-6
Σ	M32977;	VASCULAR ENDOTHELIAL GROWTH FACTOR PRECURSOR (VEGF) (VASCULAR
<u> </u>	M27281]	PERMEABILITY FACTOR) (VPF).
Σ	M35410	IGFBP-2 [INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 2
Σ	M36717	PLACENTAL RIBONUCLEASE INHIBITOR [Ribonuclease/angiogenin inhibitor RAI]
Σ	M37722;	BASIC FIBROBLAST GROWTH FACTOR RECEPTOR 1 PRECURSOR (BFGF-R) (EC
×	X66945;	2.7.1.112) (FMS-LIKE TYROSINE KINASE-2) (C-FGR) (FGFR1) (FLG) (FGFBR) (FLT2).
	M63887;	(HBGF-R-ALPHA-A1) (HBGF-R-ALPHA-A2) (HBGF-R-ALPHA-A3) + FGFR SECHETED
Σ	M63888;	FORM (M34188)
≅	M63889;M3418	
9	M34641]	- 1
	M57230	INTERLEUKIN-6 RECEPTOR BETA CHAIN [membrane glycoprotein gp130]
Σ	M57399;	PLEIOTROPHIN PRECURSOR (PTN) (HEPARIN-BINDING GROWTH-ASSOCIATED
<u>~</u>	X52946;	MOLECULE) (HB-GAM) (HEPARIN-BINDING GROWTH FACTOR 8) (HBGF-8)
<u> </u>	D90226)	(OSTEOBLAST SPECIFIC FACTOR 1) (OSF-1) (HEPARIN-BINDING NEUHITE
		OUTGROWTH PROMOTING FACTOR 1) (HBNF-1).
2	M57502	삤
2	M57765	INTERLEUKIN-11 [adipogenesis inhibitory factor]
2	M59818	GRANULOCYTE COLONY STIMULATING FACTOR RECEPTOR
<u> </u>	M59964	
2	M60278	HEPARIN-BINDING EGF-LIKE GROWTH FACTOR [DIPHTHERIA TOXIN RECEPTOR]
2	M60718	HEPATOCYTE GROWTH FACTOR PRECURSOR (SCATTER FACTOR) (SF)
		(her Allor Cell III A).
≥	M60828	FGF-7; KEHAJINOCYTE GHOWIH FACTOR PRECORSON (NGT) (FIBNOBLAS) GROWTH FACTOR-7) (HBGF-7)
2	M61176	BRAIN-DERIVED NEUROTROPHIC FACTOR
2	M62302	GDF-1 GROWTH/DIFFERENTIATION FACTOR 1]
2	M62505	CSA ANAPHYLATOXIN CHEMOTACTIC RECEPTOR
2	M65199	ENDOTHELIN-2
2	M65290	INTERLEUKIN-12 BETA CHAIN [Natural killer cell stimulatory factor, p40]
2	M65291	INTERLEUKIN-12 ALPHA CHAIN [Natural killer cell stimulatory factor, p35]
2	M67454	FASL RECEPTOR [Fas antigen, APO-1 antigen]
2	M68932	INTERLEUKIN-8 RECEPTOR (ALFA, HIGH AFFINITY)
2	M73482	NEUROMEDIN-B RECEPTOR

TABLE 8 (CONT)

	GenBank #	Gene Name
	M74178	HEPATOCYTE GROWTH FACTOR-LIKE (macrophage-stimulating protein (MS11))
	M76125	AXL (TYROSINE-PROTEIN KINASE RECEPTOR UFO)
	M92381	THYMOSIN BETA-10
	M92934	CONNECTIVE TISSUE GROWTH FACTOR
	M96956;	TDGF1 (TERATOCARCINOMA-DERIVED GROWTH FACTOR 1) (EPIDEHMAL GROWTH
	[M96955]	FACTOR-LINE CHILD PROTEIN CAT) (CAILTON GROWTH FACTOR 2) (EPIDERMAL GROWTH)
		FACTOR-LIKE CRIPTO PROTEIN CR3) (CRIPTO-3 GROWTH
	S59184	TYROSINE-PROTEIN KINASE RYK (RYK receptor-like lyrosine kinase)
	U01134;	VASCULAR ENDOTHELIAL GROWTH FACTOR RECEPTOR 1 PRECURSOR (EC
	[X51602]	2.7.1.12) (VEGFH-1) (I YHOSINE-PHOTEIN KINASE HECETOR FLIT (VETT) (VEGFH-1) (I YHOSINE-PHOTEIN KINASE
	U02687	FL CYTOKINE RECEPTOR PRECURSOR (EC.Z.1.112) (1100SINE-FROTEIN NINGSE RECEPTOR ELT3) (STEM CELL TYROSINE KINASE 1) (STK-1) (CD135 ANTIGEN).
	1103187	INTERLEUKIN-12 RECEPTOR
	U03882	C-C CHEMOKINE RECEPTOR [Monocyte chemoattractant protein 1 receptor (MCP-1RA)
		alternatively spliced
	U03905	C-C CHEMOKINE RECEPTOR [Monocyte chemoattractant protein 1 receptor (MCP-1RB)
		alternatively spliced
	U04806;	SL CYTOKINE PRECURSOR (FLT3/FLK2 LIGAND).
	[03858]	T District to the second secon
	U10117	ENDOTHELIAL-MONOCYTE ACTIVATING POLYPEPTIDE II
	U11814;	FIBROBLAST GROWTH FACTOR RECEPTOR 2 PECURSOR (FCFR-2)
	[M80634;	(KERATINOCYTE GROWTH FACTOR RECEPTOH) (FGFH2) (BER) (BFH-1) (NSAM-1) + N-
	X52832;	SAM, K-SAM-III; K-SAM-IV
	M35718;	
	M87771;	
	U14407	INTERLEUKIN-15
	U14722	ACTIVIN TYPE I RECEPTOR
	U43142	VASCULAR ENDOTHELIAL GROWTH FACTOR C PRECURSOR (VEGF-C) (VASCULAR ENDOTHELIAL GROWTH FACTOR RELATED PROTEIN) (VRP) (FLT4 LIGAND).
	Y06182	C.KIT PROTO-ONCOGENE Imast/stem cell growth factor receptor
	X06233	1=
	X06234	CALGRANULIN (A) [MRP-8 (calcium binding protein in macrophages, MIF-related)]
	X06374	Ĕĺ
:	X13967	LEUKAEMIA INHIBITORY FACTOR [cholinergic differentiation factor]
	X17543	INTERLEUKIN-9
	X17648	GRANULOCYTE-MACROPHAGE COLONY-STIMULATING FACTOR RECEPTOR ALPRA
		the first control of the first

TABLE 8 (CONT)

General Manne Factor Precursor (HBGF-1) (Acidic Fibrobia General Manne Factor General Manne Gene			
	-	Genbank #	gene Name
	- •	X51943;	HEPARIN-BINDING GROWTH FACTOR 1 PRECURSOR (HBGF-1) (ACIDIC TIBROLAS)
		[M13361;	GROWTH FACTOR!) (AFGF) (BETA-ENDOTHELIAL CELL GROWTH FACTOR) (ECGP.
		X65/78	BETA).
		X53655;	NT-3 (NEUROTROPHIN-3 PHECOHSOH) (NEUROTHIC FACTOH) (NEUVE
		[M37763]	
		X53799	MACROPHAGE INFLAMMATORY PROTEIN-2-ALPHA [MIPZalpha]
		X54936	PLACENTA GROWTH FACTORS 1 AND 2 PRECURSOR (PLGF-1 / PLGF-2).
		X59770	INTERLEUKIN-1 RECEPTOR TYPE II
		X60592	CDW40; NERVE GROWTH FACTOR RECEPTOR-RELATED B-LYMPHOCYTE
		X73304	CORTICOTROPIN RELEASING FACTOR RECEPTOR
		X78686	NEITROPHII ACTIVATING PROTEIN ENA-78
		6265LX	OX40 LIGAND [0034]
		Y00787	8
		270519	FAS/APO 1
		D17517	
		J03241	TRANSFORMING GROWTH FACTOR (BETA 3)
		J03634	INHIBIN BETA (A CHAIN) [activin A, activin AB alpha polypeptide; erythroid differentiation
		L32976	PROTEIN KINASE MLK-3 [MIXED LINEAGE KINASE 1]
		L35233	AUTOCRINE MOTILITY FACTOR RECEPTOR [AMFR]
		M31213;	PROTO ONCOGENE TYROSINE-PROTEIN KINASE RECEPTOR RET PRECURSON (EC
		[M57464]	2.7.1.112) (C-RET). (Papillary thyroid carcinoma-encoded protein)
		M95489	ĺ
		U05875	
			factor-1 (AF-1)]
		U15979;	DELTA-LIKE PROTEIN PRECURSOR (CONTAINS: FETAL ANTIGEN 1) (FA1) (DLK) +
1 0 0 0 0		[212172]	ADRENAL SPECIFIC 30kg PHOLEIN GB: X1/544
,		X03541	HIGH AFFINITY NERVE GROWTH FACTOR RECEPTOH PHECUHSON (EC 2.7.1.112) [/TRK1_TRANSFORMING TYROSINE KINASE PROTEIN] (P140-TRKA) + trk-T3 (P68 TRK)
3 0 1 2 2			T3 ONCOPROTEIN)
		X15218	SKI ONCOGENE
		X15219	
160		X74979	
: [6]		A06925	RELAXIN H2
-: m		D10232	RENIN-BINDING PROTEIN
		M13981	INHIBIN ALPHA CHAIN
		M31159;	IGFBP3 (GROWTH HORMONE-DEPENDENT INSULIN-LIKE GROWTH FACTOR-BINDING
		[M35878]	PROTEIN
		006863	FOLLISTATIN-RELATED PHOTEIN
		S85655	PROHIBITIN

TABLE 8 (CONT)

4 4440	Control Name
Geilbann #	Actio Name
D38122:	FAS ANTIGEN LIGAND (APOPTOSIS ANTIGEN LIGAND) (APTL) (APTILGT) (FASC).
[1008137]	
L11015	LYMPHOTOXIN-BETA
057059	FAS ANTIGEN LIGAND [TMF-related apoptosis inducing ligand TRAIL; Apo-2 ligand]
X14454	INTERFERON REGULATORY FACTOR [Interferon regulatory factor 1]
Y09392;	WSL-LR, WSL-S2 + TRAMP (Apo-3) (DDR3)
[U75380;U7461	
1; 083597	
M27544	INSULIN-LIKE GROWTH FACTOR IA
M86528	NEUROTROPHIN-4
M86528;	NT-4 (NT-5) + NT-6
S41541;	
[S41540;	
S41522	- 1
U14187	- 1
U14188	RECEPTOR TYROSINE KINASE LIGAND LERK-4 (EPLG4)
U32659	INTERLEUKIN-17
U33635	HIGH AFFINITY NERVE GROWTH FACTOR RECEPTOR [colon carcinoma kinase-4
	(CCK4)
U68162	THROMBOPOEITIN RECEPTOR
A25270	IFN-GAMMA ANTAGONIST CYTOKINE
A03911	NEURITE PROMOTING FACTOR(NEXIN), glia derived
D49493	BONE MORPHOGENETIC PROTEIN 3B
D49742;	HGF ACTIVATOR LIKE
[583182]	
L17075	TGF-b superfamily receptor type I (ALK-1) (SRK3)
L03840	FGFR4
L19063	GDNF
L37882	frizzled
L20861	Wnt-5a
M62403	IGFBP4
M65062	IGFBP5
M73980	Notch1
M97016	BONE MORPHOGENETIC PROTEIN 8 (OSTEOGENIC PROTEIN 2)
M99437	notch group protein (N)
U43318	lrizzled 5
X07876	WNT2 OR IRP
A26792	CNTF, ISOFORM B AND C
L42379	BPGF-1
271621	Wnt-13
M21626	T CELL RECEPTOR VARIABLE REGION

TABLE 8 (CONT)

7 7 2 2	October 18 and 1
Genbank #	
M25639	MIF
U82169	(rizzled homolog (FZD3)
U83508	angiopoietin-1
U84401	smoothened
U90875	cytotoxic ligand TRAIL receptor
U95299	Notch4
X91940	WNT-8B
X97057	WNT-10B
AF003521	Jagged 2
AF028593	Jagged 1
U77493	Notch2
U94352	manic fringe
U94354	lunatic fringe
M27968	FGF2; HEPARIN-BINDING GROWTH FACTOR 2 PRECURSOR (PROSTATROPIN). (HBGF4
138518	sonir hadaahaa (SHH)
M60314	BONE MORPHOGENETIC PROTEIN 5
MED315	BONE MORPHOGENETIC PROTEIN 6
M60316	BONE MORPHOGENETIC PROTEIN 7 (OSTEOGENIC PROTEIN 1)
D13365:	GROWTH INHIBITORY FACTOR (METALLOTHIONEIN-III)
 [M93311]	
U46010	HGF AGONIST/ANTAGOINST
L36034	SDF1A (pre-B cell stimulating factor homologue)
M15530	BCGF1 (B-cell growth factor)
M58051;	FGFR3 (FLG-2)
[X58255]	
M77227	COMPETITIVE HEPATOCYTE GROWTH FACTOR ANTAGONIST. AN ALTERNATIVE
	TRANSCRIPT OF THE REPATOCYTE GROWTH FACTOR PRECURSOR (SCALLER FACTOR) (SF) (HEPATOPOEITIN A)
U24163;	Irizzled-related FrzB (Fritz) (frezzled (fre))
[U91903; U68057]	
U28811;	CYSTEINE-RICH FIBROBLAST GROWTH FACTOR RECEPTOR [Golgi membrane
[U64791]	sialoglycoprotein MG160 (GLG1))
U48801;	VASCULAR ENDOTHELIAL GROWTH FACTOR B PRECURSOR (VEGF-B) + VEGF
[U43368]	RELATED FACTOR ISOFORM VIF 186 PRECURSOR
X02492	LEUKOCYTE INTERFERON-INDUCIBLE PEPTIDE
X85960	trk-T3 (P68 TRK-T3 ONCOPROTEIN)
X14445	FGF-3 : INT-2 PROTO-ONCOGENE PROTEIN PRECURSOR (FIBROBLAST GROWTH FACTOR-31/HBGF-3)
M37825	FGF-5; FIBROBLAST GROWTH FACTOR-5 PRECURSOR (HBGF-5).

TABLE 8 (CONT)

GenBank # Gene Name	Gene Name apoptosis-related protein TFAR15 (TFAR15) extracellular matrix metalloproteinase inducer EMMPRIN extracellular matrix metalloproteinase inducer EMMPRIN (LERK-1) (MINEDIATE EARLY RESPONSE PROTEIN B61) (TUMOR NECROSIS FACTOR, (LERK-1) (UMCEDIATE EARLY RESPONSE PROTEIN B61) (TUMOR NECROSIS FACTOR, ALPHA-INDUCEDIATE EARLY RESPONSE PROTEIN B61) (TYROSINE-PROTEIN EPHRIN TYPE-8 RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN EPHRIN-B1 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) (LERK-2) (ELK LIGAND PRECURSOR) (ELK-L). CD278P (Siva) EPHRIN-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 3) (LERK-7) (AL-1). EPHRIN-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) (LERK-R) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND BOTEIN) (LERK-B) (EPH-RELATED RECEPTOR FRANSMEMBRANE LIGAND BOTEIN) (LERK-B) (EPH-RELATED RECEPTOR SPRECURSOR)
	is-related protein TFAR15 (TFAR15) lular matrix metalloproteinase inducer EMMPRIN 1-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 1) 1-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 1) 1 (IMMEDIATE BARCY RESPONSE PROTEIN 861) (TUMOR NECROSIS FACTOR, INDUCED PROTEIN 4). 1 TYPE-8 RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN ERECEPTOR HTK). 1-81 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) 2) (ELK LIGAND PRECURSOR) (ELK-L). 2) (ELK LIGAND PRECURSOR) (ELK-L). 4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4 (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). 5) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND BAPTEIN ELY ALCEPTOR ELY 1 ACED HONDOR CON KINASE. 1 (1907) (190
	ular matrix metalloproteinase inducer EMMPRIN 1-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 1) 1-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 1) 1-41 PRECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN 4). 1-41 PRECEPTOR HTK). 2-4-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) 2-4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 3-4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-4-5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-4-5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-4-6 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-6 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-6 PRECEPTOR SPRECURSOR (EC 2.7.1.112) (TYPE-A RECEPTOR SPRECEPTOR PRECEPTOR PROTEIN A PRECEPTOR PROTEIN A PR
	1-41 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 1) (IMMEDIATE EARLY RESPONSE PROTEIN B61) (TUMOR NECROSIS FACTOR, INDUCED PROTEIN 4). 1 TYPE-B RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN 2). 1 TYPE-B RECEPTOR HTK). 1-81 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2). 1 (SIVA) 1-82 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7). 1 (AL-1). 1 (AL-1). 1 (APPE-A RECEPTOR TRANSMEMBANE LIGAND ELK-L3). 1 (EPH-RELATED RECEPTOR TRANSMEMBANE LIGAND BROTEIN. 1 (PEP-A RECEPTOR S PRECURSOR (EC 2.7.1.112) (TYPOSINE-PROTEIN.
	1) (IMMEDIATE EARLY RESPONSE PROTEIN B61) (TUMOR NECROSIS FACTOR, INDUCED PROTEIN 4). 1 TYPE-B RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN 4.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1.1
	INDUCED PHOLEIN 4). INDUCED PHOLEIN 4). I TYPE-B RECEPTOR 4 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN ERCEPTOR HTK). 4-B1 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) 2) (ELK LIGAND PRECURSOR) (ELK-L). 4-A5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 7) (AL-1). 4-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 9) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). 17 YPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN).
	1 TYPE-B RECEPTOR 4 PRECURSOR (EC 2.7.1.12) (1740 SINE-PROTEIN 1 RECEPTOR HTK). 4-81 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) 2) (ELK LIGAND PRECURSOR) (EL K-L). 4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 7) (AL-1). 4-83 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 9) (EPH-RELATED RECEPTOR TRANSMEMBANE LIGAND ELK-L3). 17 YPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN.
898888	4-B1 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 2) (ELK LIGAND PRECURSOR) (ELK-L). 9 (Siva) 4-A5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 4-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 9) (EPH-RELATED RECEPTOR TRANSMEMBANE LIGAND ELK-L3). 9) (EPH-RELATED RECEPTOR TRANSMEMBANE LIGAND ELK-L3). 17 PPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN.
8988	9 (Siva) 4-45 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 4-87 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 4-83 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 6-94 RECEPTOR TRANSMEMBRANE LIGAND ELK-L3) 7 (TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYPOSINE-PROTEIN
8988	4-A5 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 7) 7) (AL-1). 4-83 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) 8) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). 7) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). 7) (TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN. 7) PROCESTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN.
	7) (AL-1). 4-83 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) B) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). A TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN
	4-B3 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 8) B) (EPH-RELATED RECEPTOR TRANSMEMBRANE LIGAND ELK-L3). A TYPE-A RECEPTOR 5 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN.
	1 (TYPE A RECEIONS SPECURSOR (EC 27.1.12) (TYPOSINE) 1 TYPE A RECEIONS S PRECURSOR (EC 27.1.112) (TYPOSINE) 1 TYPE SINGER SPECURSOR (EC 27.1.112) (TYPOSINE)
	TOTAL STATE OF THE PROPERTY OF VINASE 11 (RECEDED BROTEIN.
	INFERIOR FIRST
	eceptor 5 (DR5)
	secreted apoptosis related protein 1
	secreted apoptosis related protein 3 (SARP3)
	EPHRIN B2 PRECURSOR (EPH-RELATED RECEPTOR TYROSINE KINASE LIGAND 5)
	INTEDITOR CHANNEL THE CONTRACTOR AND A DECEDITOR AND A DECEDIT
	TEUNIN I DESCRIPTION AND FOUND FOR 9 7 4 449) (TYDORINE DECTEIN
	EPHRIN 17PE-8 RECEPTOR 1 PRECORSON (EC 27.1.112) (1700SINE-1701EIN KINASE RECEPTOR EPH-2) (NET).
	EPHRIN TYPE-B RECEPTOR 2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN EPH-
	TYROSÍNE-PROTEIN KINASE HCK (EC 2.7.1.112) (PS9-HCK AND P60-HCK) (HEMOPOIETIC CELL KINASE).
	EPHRIN TYPE-A RECEPTOR 2 PRECURSOR (EC 2.7.1.112) (TYROSINE-PROTEIN
	KINASE RECEPTOR EUN (EPTITIBELIAL CELL NIMASE).
	GELACTIVATION OF THE CONTROL OF THE
	IGFBP COMPLEX ACID LABILE CHAIN
	FGF-8; ANDROGEN-INDUCED GROWTH FACTOR PRECURSOR (AIGF) (HBGF-8) (FIBROBLAST GROWTH FACTOR-8)
U41745 PDGF assoc. protein	assoc. protein
U43148 patched homolog	patched homolog (PTC)
J02958 MET	

TABLE 8 (CONT)

Cond Mano	Series Name
Genoans #	
U66197	FHF:1
X52599	BETA NGF
X52773	retinoic acid receptor alpha [RETINOIC ACID RECEPTOR RXR-ALPHA (RXRA)]
	FGF-6: FIBROBLAST GROWTH FACTOR-8 PRECURSOR (HBGF-6) (HST-2):
	FAU

Cell Cycle Array

In the cell cycle array according to the subject invention, all of the unique polynucleotide probe compositions correspond to genes that are associated with the life cycle of a cell. In a specific cell cycle array of interest, the spots are as provided in Table 9.

TABLE 9

	GonBank #	Gone Name
	740000.0140041	MOM2-A (GB: U33199) + MDM2-C (GB: U33199) + MDM2-C (GB: U33190) +
	Z12020; [M92424]	033201)
:	M14694; [M14695]	
	U18422	DP2 (Hundp2), dimerization partner of E2F
		DUAL SPECIFICITY MITOGEN ACTIVATED PROTEIN KINASE KINASE 1 (EC 2.7.1) (MAP
	L05624	NINASE KINASE 1) (MAPKK 1) (EHK ACTIVATOH KINASE 1) (MAPK/EHK KINASE 1) (MEK1).
	L07540	ACTIVATOR 1 36 KD SUBUNIT (REPLICATION FACTOR C 36 KD SUBUNIT) (RFC36)
	L07541	ACTIVATOR 1 38 KD SUBUNIT (REPLICATION FACTOR C 38 KD SUBUNIT) (RFC38)
	00000	CELL DIVISION PROTEIN KINASE 7 (EC 2.7.1) (CDK-ACTIVATING KINASE) (CAK) (39
	L20320	KD PHOTEIN KINASE) (P39 MOTS) (STK1) (CAK1).
	L29511; [M96995]	GROWTH FACTOR RECEPTOR-BOUND PROTEIN 2 (GRB2 ADAPTOR PROTEIN) (ASH PROTEIN).
	L33264	CDC2-RELATED KINASE PISSLRE
	M63488	REPLICATION PROTEIN A 70 KD DNA-BINDING SUBUNIT (RP-A) (RF-A) (REPLICATION FACTOR-A PROTEIN 1) (SINGLE STRANDED DNA-BINDING PROTEIN)
	M74524	HHR6A (YEAST RAD6 HOMOLOG) (UBIOITIN-CONJUGATING ENZYME) (JIRCA)
	M87338	ACTIVATOR 1 40 KD SUBUNT (REPLICATION FACTOR C 40 KD SUBUNIT) (RFC40)
	M87339	ACTIVATOR 1 37 KD SUBUNIT (REPLICATION FACTOR C 37 KD SUBUNIT) (REC37)
		OVCI IN DEPENDENT KINASE INHIBITOR 1 (MEI ANOMA DIEEEBENTIATION
		ASSOCIATED PROTEIN 6) (MDA-6) (P21) (CDK-INTERACTING PROTEIN 1) (CIP1)
	U09579; [L25610]	(WAF1) (CDKN1A) (CDKN1) (SD11) (PIC1) (CAP20)
	M68520	CELL DIVISION PROTEIN KINASE 2 (EC 2.7.1) (P33 PROTEIN KINASE)
	M81933	odc25A; M-PHASE INDUCER PHOSPHATASE 1 (EC 3.1.3.48)
` <u>;</u>	M92287	CYCLIN D3
	M96684	TRANSCRIPTIONAL ACTIVATOR PROTEIN PUR-ALPHA
	X51688	CYCLINA
	X03484	RAF ONCOGENE
	X59798; [M64349]	CYCLIN D1 (CYCLIN PRAD1) (BCL-1 ONCOGENE)
	D13639 [M90813]	ı
	HT3218 [K00065]	SUPEROXIDE DISMUTASE (Superoxide dismutase 1 (Cu/Zn))
		UV EXCISION REPAIR PROTEIN PROTEIN PAD23 (xeroderma pigmentosum group C
	D21235	repair complementing protein HHR23A)
	U11791 [U12685]	CYCLIN H
	L26318	STRESS-ACTIVATED PROTEIN KINASE JNK1 (EC 2.7.1) (C-JUN N-TERMINAL KINASE 1) (JNK-46)
	1 27211	CYCLIN-DEPENDENT KINASE 4 INHIBITOR A (CDK4) (P16-INK4) (P16-INK4A)
	1-6/611	I(MULTITLE TUMON SOFTNESSON I) (MIST). (CUNNIZA)

TABLE 9 (CONT)

	2 4 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5 5	AUTOCEN ACTIVATED BEOTEIN KINACE D38 IEC 37 4 3 MAD KINACE D38 ICVTOKINE
نِ		MILOGEN-ACTIVALED FROTEIN NINASE F36 (EC. 2.7.1.7) (NAT NINASE F36) (CTTONINE SUPPRESSIVE ANT-INITAMMATORY DRUG BINDING PROTEIN) (CSAID BINDING DEOTEIN) (CSB) (MAY INTERACTING PROTEIN 9) (MAP KINASE MY)9)
78	113228	N-mark
2	113560	
2	M15400	Retinoblastoma susceptibility (RB1 retinoblastoma-assoc)
	M25753	CYCLIN B1 G2/MITOTIC-SPECIFIC
		GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD45 (DNA-DAMAGE
2	M60974	INDUCIBLE IMANSCHIPT 1) (DUITT).
2	M73812	CYCLINE
S	\$40706 [\$62138]	GROWTH ARREST AND DNA-DAMAGE-INDUCIBLE PROTEIN GADD153 (DNA-DAMAGE INDUCIBLE PROTEIN) (CHOP).
2	U40343; [U20498]	CYCLIN-DEPENDENT KINASE 4 INHIBITOR D (P19-INK4D).
3	U47413 [L49504]	CYCLIN G1
2	U47414 [L49506]	CYCLIN G2
		EXTRACELLULAR SIGNAL-REGULATED KINASE 1 (EC 2.7.1) (ERK1) (INSULIN-
×	X60188	SHIMOLALED MARE MINASE) (MAF MINASE I) (MARIN I) (FAREINI) (EN E) (FAREINIA) (MICROTUBULE-ASSOCIATED PROTEIN-2 KINASE).
		EXTRACELLULAR SIGNAL-REGULATED KINASE 3 (EC 2.7.1.) (FRK3) (MAP KINASE
<u>×</u>	X80692	ISOFORM P97) (P97-MAPK).
		STRESS-ACTIVATED PROTEIN KINASE JNK2 (EC 2.7.1) (C-JUN N-TERMINAL KINASE
-	131931	2) (JIN -3.2).
	U34819; [U07620]	STRESS-ACTIVATED PROTEIN KINASE JNK3 (EC 2.7.1) (C.JUN N-TERMINAL KINASE 3) (JNK3) (MAP KINASE P49 3F12).
	29216	CLK-2
	29220	CLK-3
<u>L.</u>	29222	CLK-1
	U10564	WEE1-LIKE PROTEIN KINASE (EC 2.7.1.112) (Wee1Hu)
		CYCLIN-DEPENDENT KINASE INHIBITOR 1C (CYCLIN-DEPENDENT KINASE INHIBITOR
	U22398	P57) (P57KIP2)
J	J33841	ATAXIA TELANGIECTASIA (ATM)
	U39657	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 6 (EC 2.7.1) (MAPI KINASE KINASE 6) (MAPKK 6) (MAPK/ERK KINASE 6) (SAPKK3)
	M81934; [S78187]	cdc25B; M-PHASE INDUCER PHOSPHATASE 2 (EC 3.1.3.48). (CDC25Hu2)
	112075-11368441	CYCLIN-DEPENDENT KINASE 4 INHIBITOR B (P14-INK4B) (P15-INK4B) (MULTIPLE
	X74262	RBA/p48
	(85133	RBQ1 retinoplastoma binding protein
<u> </u>	X85753	CELL DIVISION PROTEIN KINASE 8 (EC 2.7.1.) (PROTEIN KINASE K35).

TABLE 9 (CONT)

		ConBoot #	Cons Name
		20000	
		L13698	GROWTH-ARREST-SPECIFIC PROTEIN 1 (GAS-1).
		D63878	NEDDS PROTEIN HOMOLOG.
		L23959	E2F-related transcription factor (DP-1)
		L25676	SERINE/THREONINE PROTEIN KINASE PITALRE
		M14505	CELL DIVISION PROTEIN KINASE 4 (EC 2.7.1) (PSK-J3)
		M29039	Jun B TRANSACTIVATOR
		M34065	cdc25C; M-PHASE INDUCER PHOSPHATASE 3 (EC 3.1.3.48).
		M35543; [M57298]	cdc42 homolog (G25K) [brain isoform + placental isoform]
		L22005	UBIQUITIN-CONJUGATING ENZYME E2-CDC34
		M95712	raf,b-
	•	872008	CDC10 PROTEIN HOMOLOG
		U15642	E2F-5
		U24152	SERINETHREONINE-PROTEIN KINASE PAK-ALPHA (EC 2.7.1) (P65-PAK) (P21- ACTIVATED KINASE) (ALPHA-PAK)
		U24153	p21-activated protein kinase (Pak2)
		1125278	EXTRACELLULAR SIGNAL-REGULATED KINASE 5 (EC 2.7.1) (ERK5) (ERK4) (BMK1 KINASE)
			CYCLIN-DEPENDENT KINASE S ACTIVATOR ISOFORM P391 PRECURSOR (CDKS
		U34051	ACTIVATOR) (P39I).
		1153442	MITOGEN-ACTIVATED PROTEIN KINASE P38 BETA (EC 2.7.1) (MAP KINASE P38
		1 24076	SCHING SECONIATED BOTTEM (EDAD)
		L340/3	PABEL-DATAMISIN ASSOCIATED PROTEIN (PRAP)
,		X05360	CELL DIVISION CONTROL PROTEIN 2 HOMOLOG (EC 2.7.1) (P34 PROTEIN KINASE) (CYCLIN-DEPENDENT KINASE 1) (CDK1)
,		L40027	glycogen synthase kinase 3
		X59727	EXTRACELLULAH SIGNAL-REGULATED KINASE 4 (EC 2.7.1) (ERK4) (MAP KINASE ISOFORM P63) (P63-MAPK).
		X66360	SERINE/THREONINE-PROTEIN KINASE PCTAIRE-2
		X66362	SERINE/THREONINE PROTEIN KINASE PCTAIRE-3
,		X66363	SERINE/THREONINE-PROTEIN KINASE PCTAIRE-1
		X66364	CELL DIVISION PROTEIN KINASE 5 (EC 2.7.1) (TAU PROTEIN KINASE II CATALYTIC SUBUNIT) (TPKII CATALYTIC SUBUNIT) (KINASE PSSALRE).
		X66365	CELL DIVISION PROTEIN KINASE 6 (EC 2.7.1) (KINASE PLSTIRE)
		X74594	RB2/p130
		X79483	EXTRACELLULAR SIGNAL-REGULATED KINASE 6 (EC 2.7.1) (ERK6) (ERK5)

TABLE 9 (CONT)

GenBank #	Gene Name
X80343	CYCLIN-DEPENDENT KINASE 5 ACTIVATOR PRECURSOR (CDK5 ACTIVATOR) (TAU PROTEIN KINASE II 23 KD SUBUNIT) (TPKII REGULATORY SUBUNIT) (P23) (P25) (P35).
X85134	RBO-3
M15796; [J04718]	PCNA (CYCLIN)
AF001954	growth inhibitor p33ING1 (ING1)
AF007111	MDM2-like p53-binding protein (MDMX)
D89667	C-myc binding protein
U66469	p53-dependent cell growth regulator CGR19
U77949	CDC6-RELATED PROTEIN
U78876	MEK KINASE 3
Y11416	p73, a monoalletically expressed p53-related protein
Y10479	E2F-3
U02570	CDC42 GTPase-activating protein
	DUAL SPECIFICITY MITOGEN-ACTIVATED PROTEIN KINASE KINASE 2 (EC 2.7.1) IMAP KINASE KINASE 2) (MAPKK 2) (ERK ACTIVATOR KINASE 2) (MAPKERK KINASE
L11285	2) (MEK2).
M63167	Aki1 (rac protein kinase alpha, protein kinase B, c-Aki)
S57153; S57160	RBP1(RETINOBLASTOMA-BINDING PROTEIN)
U23435; U31089	Abl interactor 2 (Abi-2) + Abl binding protein 3 (AbIBP3) [ArgBPIB]
M29870: [M31467]	RAS-RELATED C3 BOTULINUM TOXIN SUBSTRATE 1 (P21-RAC1) (RAS-LIKE PROTEIN TC25)
1 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2 2	Ent. 1 De hinding acciding
M96577	EZF-1 PRD-UNDUR PIDENT PROTECTION ACTIVATED BENTEIN KINASE KINASE 6 (FC 9 7 1.) (MAP
U25265	DUAL SPECIFICITY MITOGEN ACTIVATED PROTEIN NIVASE NIVASE STORY (PORTION OF STORY) (MAPKK 5) (MAPKK 5) (MAPKK 5) (MAPKK 5).
X66357	CELL DIVISION PROTEIN KINASE 3 (EC 2.7.1).
M74091	CYCLIN C G1/S-SPECIFIC
M80629	CDC2-RELATED PROTEIN KINASE CHED
S66431	RBP2 retinoblastoma binding protein
100001	CDC27HS PROTEIN
U01038	SERINE/THREONINE-PROTEIN KINASE PLK (EC 2.7.1) (PLK-1) (STPK13)
D50310	CYCLIN I
U18291	CDC16HS.
U63131	CDC37 HOMOLOG.
U69276	GRB-IR / GRB10
X66358	SEBINE THREONINE - PROTEIN KINASE KKIALBE

Other Representative Arrays

In a neuroarray according to the subject invention, all of the unique polynucleotide probe compositions will correspond to genes that are expressed in brain related tissues. Genes that are represented on the array are key genes, by which is meant that they have been reported to play primary roles in a variety of different biological processes in brain tissues. Genes of interest that may be represented on the array include: ion channel/transport proteins; receptors; cell cycle regulators; stress response proteins; apoptosis proteins; signal transduction proteins; transcriptional factors; growth factors/interleukins/hormones; oncogenes and tumor suppressors; cell surface/adhesion proteins; DNA synthesis/repair/recombination genes; and metabolic pathway enzymes.

In certain embodiments, of particular interest is an array having the following types of genes represented on its surface: nuclear proteins; endoplasmic reticulum proteins; golgi complex proteins; endosomal proteins; lysosomal proteins; peroxisomal proteins; mitochondrial proteins; cytoplasmic proteins; cytoskeletal proteins; plasma membrane proteins; post synaptic and dendritic proteins; axonal and nerve terminal proteins; secreted proteins, neuropeptides, hormones and growth factors; extracellular matrix proteins; astrocyte and oligodendroglial proteins; immune system proteins; developmentally regulated proteins; regionally regulated proteins; and disease related proteins.

Other representative arrays include: (1) rat arrays, in which each of the unique polynucleotide corresponds to a key rat gene; (2) blood arrays, in which each unique polynucleotide corresponds to a gene associated cells and tissues associated with the cardiovascular system; (3) rat stress arrays; and (4) mouse stress arrays, in which each unique polynucleotide corresponds to a gene associated with the stress response of murine cells.

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METHODS OF USING THE SUBJECT ARRAYS

The subject arrays find use in a variety of different applications in which one is interested in detecting the occurrence of one or more binding events between target nucleic acids and probes on the array and then relating the occurrence of the binding event(s) to the presence of a target(s) in a sample. In general, the device will be contacted with the sample suspected of containing the target under conditions sufficient for binding of any target

present in the sample to a complementary polynucleotide present on the array. Generally, the sample will be a fluid sample and contact will be achieved by introduction of an appropriate volume of the fluid sample onto the array surface, where introduction can be pipette, deposition, and the like.

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Generation of Labeled Target

Targets may be generated by methods known in the art. mRNA can be labeled and used directly as a target, or converted to a labeled cDNA target. Generally, such methods include the use of oligonucleotide primers. Primers that may be employed include oligo dT, random primers, e.g. random hexamers and gene specific primers.

Of particular interest in the generation of labeled target is the use of a set of a representational number of gene specific primers, as described in U.S. Patent Application No. 08/859,998, the disclosure of which is herein incorporated by reference. As the subject sets comprise a representational number of primers, the total number of different primers in any given set will be only a fraction of the total number of different or distinct RNAs in the sample, where the total number of primers in the set will generally not exceed 80 %, usually will not exceed 50 % and more usually will not 20% of the total number of distinct RNAs, usually the total number of distinct messenger RNAs (mRNAs), in the sample. Any two given RNAs in a sample will be considered distinct or different if they comprise a stretch of at least 100 nucleotides in length in which the sequence similarity is less than 98%, as measured using the FASTA algorithm at default settings. As the sets of gene specific primers comprise only a representational number of primers, with physiological sources comprising from 5,000 to 50,000 distinct RNAs, the number of different gene specific primers in the set of gene specific primers will typically range from about 20 to 10,000, usually from 50 to 2,000 and more usually from 75 to 1500.

Each of the gene specific primers of the sets described above will be of sufficient length to specifically hybridize to a distinct nucleic acid member of the sample, e.g. RNA or c DNA, where the length of the gene specific primers will usually be at least 8 nt. more usually at least 20 nt and may be as long as 25 nt or longer, but will usually not exceed 50 nt. The gene specific primers will be sufficiently specific to hybridize to complementary template sequence during the generation of labeled nucleic acids under conditions sufficient for first strand cDNA synthesis, which conditions are known by those of skill in the art. The

number of mismatches between the gene specific primer sequences and their complementary template sequences to which they hybridize during the generation of labeled nucleic acids in the subject methods will generally not exceed 20 number %, usually will not exceed 10 number % and more usually will not exceed 5 number %.

Generally, the sets of gene specific primers will comprise primers that correspond to at least 20, usually at least 50 and more usually at least 75 distinct genes as represented by distinct mRNAs in the sample, where the term "distinct" when used to describe genes is as defined above, where any two genes are considered distinct if they comprise a stretch of at least 100 nt in their RNA coding regions in which the sequence similarity does not exceed 98%, as determined using the FASTA algorithm at default settings.

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The gene specific oligonucleotide primers may be synthesized by conventional oligonucleotide chemistry methods, where the nucleotide units may be: (a) solely nucleotides comprising the heterocyclic nitrogenous bases found in naturally occurring DNA and RNA, e.g. adenine, cytosine, guanine, thymine and uracil; (b) solely nucleotide analogs which are capable of base pairing under hybridization conditions in the course of DNA synthesis such that they function as the above nucleotides found in naturally occurring DNA and RNA, where illustrative nucleotide analogs include inosine, xanthine, hypoxanthine, 1,2-diaminopurine and the like; or (c) from combinations of the nucleotides of (a) and nucleotide analogs of (b), where with primers comprising a combination of nucleotides and analogues thereof, the number of nucleotide analogues in the primers will typically be less than 25 and more typically less than 5. The gene specific primers may comprise reporter or hapten groups, usually 1 to 2, which serve to improve hybridization properties and simplify detection procedure.

Depending on the particular point at which the gene specific primers are employed in the generation of the labeled nucleic acids, e.g. during first strand cDNA synthesis or following one or more distinct amplification steps, each gene specific primer may correspond to a particular RNA by being complementary or similar, where similar usually means identical, to the particular RNA. For example, where the gene specific primers are employed in the synthesis of first strand cDNA, the gene specific primers will be complementary to regions of the RNAs to which they correspond.

Each gene specific primer can be complementary to a sequence of nucleotides which is unique in the population of nucleic acids, e.g. mRNAs, with which the primers are

contacted, or one or more of the gene specific primers in the set may be complementary to several nucleic acids in a given population, e.g. multiple mRNAs, such that the gene specific primer generates labeled nucleic acid when one or more of set of related nucleic acid species, e.g. species having a conserved region to which the primer corresponds, are present in the sample. Examples of such related nucleic acid species include those comprising: repetitive sequences, such as Alu repeats, Al repeats and the like; homologous sequences in related members of a gene-family; polyadenylation signals; splicing signals; or arbitrary but conversed sequences.

Depending on the particular nature of the labeled nucleic acid generation step of the subject methods, the gene specific primers may be modified in a variety of ways. One way the gene specific primers may be modified is to include an anchor sequence of nucleotides, where the anchor is usually located 5' of the gene specific portion of the primer and ranges in length from 10 to 50 nt in length, usually 15 to 40 nt in length. The anchor sequence may comprise a sequence of bases which serves a variety of functions, such as a sequence of bases which correspond to the sequence found in promoters for bacteriophage RNA polymerase, e.g. T7 polymerase, T3 polymerase, SP6 polymerase, and the like; arbitrary sequences which can serve as subsequent primer binding sites; and the like.

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Turning now to the methods employing the above sets of gene specific primers, the first step in the subject methods is to obtain a sample of nucleic acids, usually RNAs, from a physiological source, usually a plurality of physiological sources, where the term plurality is used to refer to 2 or more distinct physiological sources. The physiological source of RNAs will typically be eukaryotic, with physiological sources of interest including sources derived single celled organisms such as yeast and multicellular organisms, including plants and animals, particularly mammals, where the physiological sources from multicellular organisms may be derived from particular organs or tissues of the multicellular organism, or from isolated cells derived therefrom. Thus, the physiological sources may be different cells from different organisms of the same species, e.g. cells derived from different humans, or cells derived from the same human (or identical twins) such that the cells share a common genome, where such cells will usually be from different tissue types, including normal and diseased tissue types, e.g. neoplastic, cell types. In obtaining the sample of RNAs to be analyzed from the physiological source from which it is derived, the physiological source may be subjected to a number of different processing steps, where such processing steps

might include tissue homogenation, nucleic acid extraction and the like, where such processing steps are known to the those of skill in the art. Methods of isolating RNA from cells, tissues, organs or whole organisms are known to those of skill in the art and are described in Maniatis *et al.*, Molecular Cloning: A Laboratory Manual (Cold Spring Harbor Press)(1989).

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The next step in the subject methods is the generation of labeled nucleic acids representative of the nucleic acid, usually RNA, profile of the physiological source. As mentioned above, a set of gene specific primers is used to generate the labeled nucleic acids from the sample of RNAs, where the labeled nucleic acids generated in this step may serve as "target" in subsequent assays in which the differences in the RNA profiles of at least two sources are analyzed. As used herein, the term "target" refers to single stranded RNA, single stranded DNA and double stranded DNA, where the target is generally greater than 50 nt in length.

The set of primers may be used either in first strand cDNA synthesis or following one or more amplification steps. Furthermore, the actual synthesis of the labeled nucleic acids may be at the same step during which the sets of gene specific primers are employed, or the synthesis of the labeled nucleic acids may be one more steps subsequent to the step in which the sets of gene specific primers are employed.

In a first embodiment of the invention, the set of gene specific primers is used to generate labeled first strand cDNA, where the labeled first strand cDNA is representative of the RNA profile of the physiological source being assayed. The labeled first strand cDNA is prepared by contacting the RNA sample with the primer set and requisite reagents under conditions sufficient for reverse transcription of the RNA template in the sample. Requisite reagents contacted with the primers and RNAs are known to those of skill in the art and will generally include at least an enzyme having reverse transcriptase activity and dNTPs in an appropriate buffer medium.

A variety of enzymes, usually DNA polymerases, possessing reverse transcriptase activity can be used for the first strand cDNA synthesis step. Examples of suitable DNA polymerases include the DNA polymerases derived from organisms selected from the group consisting of a thermophilic bacteria and archaebacteria, retroviruses, yeasts, Neurosporas, Drosophilas, primates and rodents. Preferably, the DNA polymerase will be selected from the group consisting of Moloney murine leukemia virus (M-MLV) as described in United

States Patent No. 4,943,531 and M-MLV reverse transciptase lacking RNaseH activity as described in United States Patent No. 5,405,776 (the disclosures of which patents are herein incorporated by reference), human T-cell leukemia virus type I (HTLV-I), bovine leukemia virus (BLV), Rous sarcoma virus (RSV), human immunodeficiency virus (HIV) and Thermus aquaticus (Taq) or Thermus thermophilus (Tth) as described in United States Patent No. 5,322,770, the disclosure of which is herein incorporated by reference. Suitable DNA polymerases possessing reverse transcriptase activity may be isolated from an organism, obtained commercially or obtained from cells which express high levels of cloned genes encoding the polymerases by methods known to those of skill in the art, where the particular manner of obtaining the polymerase will be chosen based primarily on factors such as convenience, cost, availability and the like.

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The various dNTPs and buffer medium necessary for first strand cDNA synthesis through reverse transcription of the primed RNAs may be purchased commercially from various sources, where such sources include Clontech, Sigma, Life Technologies, Amersham, Boehringer-Mannheim. Buffer mediums suitable for first strand synthesis will usually comprise buffering agents, usually in a concentration ranging from 10 to 100 µM which typically support a pH in the range 6 to 9, such as Tris-HCl, HEPES-KOH, etc.; salts containing monovalent ions, such as KCl, NaCl, etc., at concentrations ranging from 0-200 mM: salts containing divalent cations like MgCl₂, Mg(OAc) etc, at concentrations usually ranging from 1 to 10 mM; and additional reagents such as reducing agents, e.g. DDT, detergents, albumin and the like. The conditions of the reagent mixture will be selected to promote efficient first strand synthesis. Typically the set of primers will first be combined with the RNA sample at an elevated temperature, usually ranging from 50 to 95 °C, followed by a reduction in temperature to a range between about 0 to 60°C, to ensure specific annealing of the primers to their corresponding RNAs in the sample. Following this annealing step, the primed RNAs are then combined with dNTPs and reverse transcriptase under conditions sufficient to promote reverse transcription and first strand cDNA synthesis of the primed RNAs. By using appropriate types of reagents, all of the reagents can be combined at once if the activity of the polymerase can be postponed or timed to start after annealing of the primer to the RNA.

In this embodiment, one of either the gene specific primers or dNTPs, preferably the dNTPs, will be labeled such that the synthesized cDNAs are labeled. By labeled is meant

that the entities comprise a member of a signal producing system and are thus detectable, either directly or through combined action with one or more additional members of a signal producing system. Examples of directly detectable labels include isotopic and fluorescent moieties incorporated into, usually covalently bonded to, a nucleotide monomeric unit, e.g. dNTP or monomeric unit of the primer. Isotopic moieties or labels of interest include ³²P, ¹³P, ³⁵S, ¹²⁵I, and the like. Fluorescent moieties or labels of interest include coumarin and its derivatives, e.g. 7-amino-4-methylcoumarin, aminocoumarin, bodipy dyes, such as Bodipy FL, cascade blue, fluorescein and its derivatives, e.g. fluorescein isothiocyanate, Oregon green, rhodamine dyes, e.g. texas red, tetramethylrhodamine, eosins and erythrosins, cyanine dyes, e.g. Cy3 and Cy5, macrocyclic chelates of lanthanide ions, e.g. quantum dye™, fluorescent energy transfer dyes, such as thiazole orange-ethidium heterodimer, TOTAB, etc. Labels may also be members of a signal producing system that act in concert with one or more additional members of the same system to provide a detectable signal. Illustrative of such labels are members of a specific binding pair, such as ligands, e.g. biotin, fluorescein, digoxigenin, antigen, polyvalent cations, chelator groups and the like, where the members specifically bind to additional members of the signal producing system, where the additional members provide a detectable signal either directly or indirectly, e.g. antibody conjugated to a fluorescent moiety or an enzymatic moiety capable of converting a substrate to a chromogenic product, e.g. alkaline phosphatase conjugate antibody; and the like.

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In one preferred embodiment, the member of the signal producing system bound to the nucleotide is functional group capable of covalently binding to additional members of the signal producing system to generate a detectable label. Examples of such functional groups or moieties include amino, sulfhydryl, azido, isothiocyanate, sulfoxyl, and the like. The labeled target generated using such nucleotides will thus include one or more, usually a plurality of, functional moieties. For detection, the functional moieties of the modified nucleotides can be labeled by conjugation of a label to the functional moiety. A variety of suitable labels and methods for their conjugation to functional moieties are known to those of skill in the art. Examples include labeling of amino-modified cDNA by a succinimidyl ester of an appropriate dye, e.g. Alexa, Bodipy, or Cy dyes. Alternatively, label can be entrapped or bonded into structures of microscopic-sized particles. These particles can then be conjugated with the functional moieties of the target.

For each sample of RNA, one can generate labeled oligos with the same labels.

Alternatively, one can use different labels for each physiological source, which provides for additional assay configuration possibilities, as described in greater detail below.

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In a variation of the above embodiment, where desired one can generate labeled RNA instead of labeled first strand cDNA. In this embodiment, first strand cDNA synthesis is carried out in the presence of unlabeled dNTPs and unlabeled gene specific primers. However, the primers are optionally modified to comprise a promotor for an RNA polymerase, such as T7 RNA polymerase, T3 RNA polymerase, SP6 RNA polymerase, and the like. In this embodiment, following first strand cDNA synthesis, the resultant single stranded cDNA is then converted to double stranded cDNA, where the resultant double stranded cDNA comprises the anchor sequence comprising the promoter region. Conversion of the mRNA:cDNA hybrid following first strand synthesis can be carried out as described in Okayama & Berg, Mol. Cell. Biol. (1982) 2:161-170, and Gubler & Hoffman, Gene (1983) 25: 253-269, where briefly the RNA is digested with a ribonuclease, such as E.coli RNase H. followed by repair synthesis using a DNA polymerase like DNA polymerase I, etc., and E.coli DNA ligase. One may also employ the modification of this basic method described in Wu, R, ed., Methods in Enzymology (1987), vol. 153 (Academic Press). Next, the double stranded cDNA is contacted with RNA polymerase and dNTPs, including labeled dNTPs as described above, to produce linearly amplified labeled ribonucleic acids. For cDNA lacking the anchor sequence comprising a promoter region, a polymerase that does not need a promoter region but instead can initiate RNA strand synthesis randomly from cDNA; such as core fragment of E.Coli RNA polymerase, may be employed.

In another embodiment of the subject invention, the labeled nucleic acid generation step comprises one or more enzymatic amplification steps in which multiple DNA copies of the initial RNAs present in the sample are produced, from which multiple copies of the initial RNA or multiple copies of antisense RNA (aRNA) may be produced, using the polymerase chain reaction, as described in U.S. Pat. No. 4.683,195, the disclosure of which is herein incorporated by reference, in which repeated cycles of double stranded DNA denaturation, oligonucleotide primer annealing and DNA polymerase primer extension are performed, where the PCR conditions may be modified as described in U.S. Pat No. 5,436,149, the disclosure of which is herein incorporated by reference.

In one embodiment involving enzymatic amplification, the set of gene-specific primers are employed in the generation of the first strand cDNA, followed by amplification of the first strand cDNA to produce amplified numbers of labeled cDNA. In this embodiment, as a set of gene-specific primers is employed in the first strand synthesis step, only a representative proportion of the total RNA in the sample is amplified during the subsequent amplification steps.

Amplification of the first strand cDNA can be conveniently achieved by using a CAPswitchTM oligonucleotide as described in U.S. Patent Application Serial No. 08/582,562, the disclosure of which is herein incorporated by reference. Briefly, the CAPswitchTM technology uses a unique CAPswitchTM oligonucleotide in the first strand cDNA synthesis followed by PCR amplification in the second step to generate a high yield of ds cDNA. When included in the first-strand cDNA synthesis reaction mixture, the CAPswitchTM oligonucleotide serves as a short extended template. When reverse transcriptase stops at the 5' end of the mRNA template in the course of first strand cDNA synthesis it switches templates and continues DNA synthesis to the end of the CAPswitchTM oligonucleotide. The resulting ss cDNA incorporates at the 3' end, sequence which is complimentary to complete 5' end of the mRNA and the CAPswitchTM oligonucleotide sequence.

Of particular interest as the CAPswitchTM oligonucleotide are oligonucleotides having the following formula:

5'-dN1-dN2-...dNm-rN1-rN2...rNn-3'

wherein:

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dN represents a deoxyribonucleotide selected from among dAMP, dCMP, dGMP and dTMP;

m represents an integer 0 and above, preferably from 10 to 50;

rN represents a ribonucleotide selected from the group consisting of AMP, CMP, GMP and UMP, preferably GMP; and

n represents an integer 0 and above, preferably from 3 to 7.

The structure of the CAPswitch™ oligonucleotide may be modified in a number of ways, such as by replacement of 1 to 10 nucleotides with nucleotide analogs, incorporation

of terminator nucleotides, such as 3'-amino NMP, 3'-phosphate NMP and the like, or non-natural nucleotides which can improve efficiency of the template switching reaction but still retain the main function of the CAPswitchTM oligonucleotide *i.e.* CAP-depended extension of full-length cDNA by reverse transcriptase using CAPswitchTM oligonucleotide as a template.

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In using the CAPswitch™ oligonucleotide, first strand cDNA synthesis is carried out in the presence of a set of gene specific primers and a CAPswitch™ oligonucleotide, where the gene specific primers have been modified to comprise an arbitrary anchor sequence at their 5' ends. The first strand cDNA is then combined with primer sequences complementary to: (a) all or a portion of the CAPswitch™ oligonucleotide and (b) the arbitrary anchor sequence of the gene specific primers and additional PCR reagents, such as dNTPs, DNA polymerase, and the like, under conditions sufficient to amplify the first strand cDNA. Conveniently, PCR is carried out in the presence of labeled dNTPs such that the resultant, amplified cDNA is labeled and serves as the labeled or target nucleic acid. Labeled nucleic acid can also be produced by carrying out PCR in the presence of labeled primers, where either or both the CAPswitchTM oligonucleotide complementary primer and anchor sequence complementary primer may be labeled. In yet an alternative embodiment, instead of producing labeled amplified cDNA, one may generate labeled RNA from the amplified ds cDNA, e.g. by using an RNA polymerase such as E.coli RNA polymerase, or other RNA polymerases requiring promoter sequences, where such sequences may be incorporated into the arbitrary anchor sequence.

Instead of using the set of gene specific primers in the first strand cDNA synthesis step followed by subsequent amplification of only a representative fraction of the total number of distinct RNA species in the sample, one may also amplify all of the RNAs in the sample and use the set of gene specific primers to generate labeled nucleic acid following amplification. This embodiment may find use in situations where the RNA of interest to be amplified is known or postulated to be in small amounts in the sample.

In this embodiment, first strand synthesis is carried out using: (a) an oligo dT primer that usually comprises an arbitrary anchor sequence at its 5' end and (b) a CAPswitchTM oligonucleotide. During first strand synthesis the oligo(dT) anneals to the polyA tail of the mRNA in the sample and synthesis extends beyond the 3' end of the RNA to include the CAPswitchTM oligonucleotide, yielding a first strand cDNA comprising an arbitrary

sequence at its 5' end and a region complementary to the CAPswitchTM oligonucleotide at its 3' end. The length of the dT primer will typically range from 15 to 30 nts, while the arbitrary anchor sequence or portion of the primer will typically range from 15 to 25 nt in length.

Following first strand synthesis, the cDNA is amplified by combining the first strand cDNA with primers that correspond at least partially to the anchor sequence and the CAPswitchTM oligonucleotide under conditions sufficient to produce an amplified amount of the cDNA. Labeled nucleic acid is then produced by contacting the resultant amplified cDNA with a set of gene specific primers, a polymerase and dNTPs, where at least one of the gene specific primers and dNTPs are labeled.

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When employed to generate target, as described above, the gene specific primers of the sets of primers according to the subject invention are typically chosen according to a number of different criteria. In some embodiments of the invention, primers of interest for inclusion in the set include primers corresponding to genes which are typically differentially expressed in different cell types, in disease states, in response to the influence of external agents, factors or infectious agents, and the like. In other embodiments, primers of interest are primers corresponding to genes which are expected to be, or already identified as being, differentially expressed in different cell, tissue or organism types. Preferably, at least 2 different gene functional classes will be represented in the sets of gene specific primers, where the number of different functional classes of genes represented in the primer sets will generally be at least 3, and will usually be at least 5. Gene functional classes of interest include oncogenes; genes encoding tumor suppressors; genes encoding cell cycle regulators; stress response genes; genes encoding ion channel proteins; genes encoding transport proteins; genes encoding intracellular signal transduction modulator and effector factors; apoptosis related genes; DNA synthesis/recombination/repair genes; genes encoding transcription factors; genes encoding DNA-binding proteins; genes encoding receptors, including receptors for growth factors, chemokines, interleukins, interferons, hormones, neurotransmitters, cell surface antigens, cell adhesion molecules etc.; genes encoding cellcell communication proteins, such as growth factors, cytokines, chemokines, interleukins, interferons, hormones etc.; and the like. Less preferred are gene specific primers that are subject to formation of strong secondary structures with less than -10kcal/mol; comprise stretches of homopolymeric regions, usually more than 5 identical nucleotides; comprise

more than 3 repetitive sequences; have high, e.g. more than 80%, or low, e.g. less than 30%, GC content etc.

The particular genes represented in the set of gene specific primers will necessarily depend on the nature of physiological source from which the RNAs to be analyzed are derived. For analysis of RNA profiles of eukaryotic physiological sources, the genes to which the gene specific primers correspond will usually be Class II genes which are transcribed into RNAs having 5' caps, e.g. 7-methyl guanosine or 2,2,7-trimethylguanosine, where Class II genes of particular interest are those transcribed into cytoplasmic mRNA comprising a 7-methyl guanosine 5' cap and a polyA tail.

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For analysis of RNA profiles of mammalian physiological sources, of particular interest are gene specific primers corresponding to the functional gene classes listed above. For analysis of RNA profiles of human physiological sources, the gene specific primers of particular interest are the gene specific primers identified in Table 1 as SEQ ID NO:01 to SEQ ID NO:1372, of U.S. Application Serial No. 08/859,998, the disclosure of which is herein incorporated by reference, where sets of these primers will usually include at least 20 and more usually at least 50 of these specific sequences.

Particular sets of primers of interest in the subject invention are those sets of primers that include primers capable of amplifying at least a portion of the unique polynucleotides present on the arrays with which the target is to be employed. By at least a portion is meant at least about 10, usually at least about 20 and more usually at least about 25 number % (where number is the number of different unique polynucleotides on the array). For examples, sets of primers that include primers capable of amplifying at least a portion of the unique polynucleotides listed in Table 1, *supra*, are of interest. Similarly sets of primers capable of amplifying at least a portion of the unique polynucleotides listed in Tables 2 to 8, *supra*, are also of interest.

In a particularly preferred embodiment, the gene specific primers are preferably those primers that correspond to the different polynucleotide spots on the array that is used in the hybridization assay. Thus, one will preferably employ gene specific primers for each different polynucleotide that is present on the array, so that if the gene is expressed in the particular cell or tissue being analyzed, labeled target will be generated from the sample for that gene. In many embodiments in which the subject arrays are employed, the gene specific primers used to generate the target from the human cell or tissue being analyzed will have

the same sequence as the gene specific primers used to generate the polynucleotide probes present on the array. In this manner, if a particular gene present on the array is expressed in a particular sample, the appropriate target will be generated and subsequently identified.

Representative sets of primers falling within this particularly preferred embodiment include:

5	SET	DESCRIPTION	
	1	I pair of primers capable of amplifying each polynucleotide listed in Table I, <i>supra</i> , as well one set of primers capable of amplifying each of the complementary sequences of each of the polynucleotides listed in Table I.	
	2	I pair of primers capable of amplifying each polynucleotide listed in Table 2, <i>supra</i> , as well one set of primers capable of amplifying each of the complementary sequences of each of the polynucleotides listed in Table 2.	
	3	I pair of primers capable of amplifying each polynucleotide listed in Table 3, supra, as well one set of primers capable of amplifying each of the complementary sequences of each of the polynucleotides listed in Table 3.	

10 Hybridization and Detection

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As mentioned above, following preparation of the target nucleic acid from the tissue or cell of interest, the target nucleic acid is then contacted with the array under hybridization conditions, where such conditions can be adjusted, as desired, to provide for an optimum level of specificity in view of the particular assay being performed. Suitable hybridization conditions are well known to those of skill in the art and reviewed in Maniatis et al, *supra* and WO 95/21944. In analyzing the differences in the population of labeled target nucleic acids generated from two or more physiological sources using the arrays described above, each population of labeled target nucleic acids are separately contacted to identical probe arrays or together to the same array under conditions of hybridization, preferably under stringent hybridization conditions (for example, at 50°C or higher and 0.1XSSC (15 mM sodium chloride/01.5 mM sodium citrate)), such that labeled target nucleic acids hybridize to complementary probes on the substrate surface.

Where all of the target sequences comprise the same label, different arrays will be employed for each physiological source (where different could include using the same array at different times). Alternatively, where the labels of the targets are different and

distinguishable for each of the different physiological sources being assayed, the opportunity arises to use the same array at the same time for each of the different target populations. Examples of distinguishable labels are well known in the art and include: two or more different emission wavelength fluorescent dyes, like Cy3 and Cy5, two or more isotopes with different energy of emission, like ³²P and ³³P, labels which generate signals under different treatment conditions, like temperature, pH, treatment by additional chemical agents, etc., or generate signals at different time points after treatment. Using one or more enzymes for signal generation allows for the use of an even greater variety of distinguishable labels, based on different substrate specificity of enzymes (alkaline phosphatase/peroxidase).

Following hybridization, non-hybridized labeled nucleic acid is removed from the support surface, conveniently by washing, generating a pattern of hybridized nucleic acid on the substrate surface. A variety of wash solutions are known to those of skill in the art and may be used.

The resultant hybridization patterns of labeled nucleic acids may be visualized or detected in a variety of ways, with the particular manner of detection being chosen based on the particular label of the target nucleic acid, where representative detection means include scintillation counting, autoradiography, fluorescence measurement, colorimetric measurement, light emission measurement and the like.

Following detection or visualization, the hybridization patterns may be compared to identify differences between the patterns. Where arrays in which each of the different probes corresponds to a known gene are employed, any discrepancies can be related to a differential expression of a particular gene in the physiological sources being compared.

Utility

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The subject methods find use in, among other applications, differential gene expression assays. Thus, one may use the subject methods in the differential expression analysis of: (a) diseased and normal tissue, e.g. neoplastic and normal tissue, (b) different tissue or tissue types; (c) developmental stage; (d) response to external or internal stimulus; (e) response to treatment; and the like. The subject arrays therefore find use in broad scale expression screening for drug discovery and research, such as the effect of a particular active agent on the expression pattern of genes in a particular cell, where such information can be

used to reveal drug toxicity, carcinogenicity, etc., environmental monitoring, disease research and the like.

Kits

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Also provided are kits for performing analyte binding assays using the subject devices, where kits for carrying out differential gene expression analysis assays are preferred. Such kits according to the subject invention will at least comprise the subject arrays. The kits may further comprise one or more additional reagents employed in the various methods, such as primers for generating target nucleic acids, including a set of gene specific primers according to the subject invention, e.g. primer sets 1 to 9 described above, dNTPs and/or rNTPs, which may be either premixed or separate, one or more uniquely labeled dNTPs and/or rNTPs, such as biotinylated or Cy3 or Cy5 tagged dNTPs, or other post synthesis labeling reagent, such as chemically active derivatives of fluorescent dyes, enzymes, such as reverse transcriptases, DNA polymerases, and the like, various buffer mediums, e.g. hybridization and washing buffers, prefabricated probe arrays, labeled probe purification reagents and components, like spin columns, etc., signal generation and detection reagents, e.g. streptavidin-alkaline phosphatase conjugate, chemifluorescent or chemiluminescent substrate, and the like.

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The following examples are offered by way of illustration and not by way of limitation.

EXPERIMENTAL

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Example 1 - Generation of human cDNA array

686 cDNA fragments corresponding 686 different human genes were amplified from quick-clone cDNA (CLONTECH) in 686 separate test tubes using a combination of sense and antisense gene-specific primers: (Set No. 9, described *supra*). Amplification was conducted in a 100-μl volume containing 2 μl of mixture of 10 Quick-clone cDNA from placenta, brain, liver, lung, leukocytes, spleen, skeletal muscle, testis, kidney and ovary (CLONTECH), 40 mM Tricine-KOH (pH 9.2 at 22°C), 3.5 mM Mg(OAc)₂, 10 mM KOAc.

75 µg/ml BSA, 200 µM of each dATP, dGTP, dCTP and dTTP, 0.2 µM of each sense and antisense gene-specific primers and 2 µl of KlenTaq Polymerase mix. Temperature parameters of the PCR reactions were as follows: 1 min at 95°C followed by 20-35 cycles of 95°C for 15 sec and 68°C for 2 min; followed by a 10-min final extension at 68°C. PCR products were examined on 1.2% agarose/EtBr gels in 1x TBE buffer. As a DNA size marker a 1 Kb DNA Ladder was used. ds cDNA was then precipitated by addition of a half volume of 4M ammonium acetate (about 35 μl) and 3.7 volumes of 95% ethanol (about 260 ul). After vortexing, the tube was immediately centrifuged at 14,000 r.p.m. in a microcentrifuge for 20 min. The pellet was washed with 80% ethanol without vortexing, centrifuged as above for 10 min, air dried, and dissolved in 10 µl of deionized water. Yield of ds cDNA after the amplification step was about 5 µg. The ds cDNA fragments for all 686. genes were cloned into TA-cloning vector using the manufacturer's recommendations (Invitrogen) and identity of the clones was confirmed by sequence analysis. The ds cDNA inserts with the sequence corresponding 686 genes were amplified by PCR using a combination of antisense and sense gene-specific primers, as described above. The ds cDNA was denatured by adding 1 µl of 10X denaturing solution (1 M NaOH, 10 mM EDTA) and incubating at 65°C for 20 min. All cDNA probes were transferred in 384-well plate and loaded on positively charged nylon membrane (Schleher & Schull) using 384 pin tool and Biomek 2000 (Beckman) robot. The resultant array is described in Table 1.

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Example 2 - Generation of ³²P-labeled oligonucleotides during first strand cDNA synthesis

Step A. cDNA synthesis/Labeling Procedure

1 μg of polyA+RNA or total RNA was converted into ³²P-labeled first-strand cDNA as follows. A sufficient volume of master mix for all labeling reactions and 1 extra reaction was prepared as follows to ensure sufficient volume. For each 10-μl labeling reaction, the following reagents were mixed:

- 2 μl 5X First-strand buffer (250 μM Tris-HCl pH8.3; 375 mM KCl; 15 mM MgCl₂)
- 1 μl 10XdNTP mix (500 μM dGTP, 500 μM dCTP, 500 μM dTTP, 5 μM dATP)
 - 4 μ l [α 32 P]dATP (Amersham, 3000 Ci/mmol, 10 mCi/ml)
 - 1 μ1 MMLV reverse transcriptase (Amersham, 200 units/μ1)

8 μl Final volume

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Next, the following reagents were combined in a 0.5-ml PCR test tube:

1 μg (1-2 μl) polyA+RNA sample

1 μl 10x gene-specific primers mix (0.2 μM of each oligonucleotide 1D No.

2,4,6,8,10,12,.... 1372 from Table 1 of U.S. Patent Application Serial No.

08/859,998, the discosure of which is herein incorporated by reference.)

10 As a control, in separate test tube were mixed 1 μg of polyA+RNA sample with 1 μl of oligo dT primer (CDS1, 5'-d(TCTAGAATTCAGCGGCCGC(T)₃₀VN) - 3'

(where V=G or A or C; N=G or A or T or C)

For each tube, ddH₂0 was added to a final volume of 3 μl and the contents were mixed and spun briefly in a microcentrifuge. The tubes were then incubated in a preheated PCR thermocycler at 70°C for 2 min. The temperature in thermocycle was reduced down to 50°C and the tube contents were incubated for 2 min. 8 μl of master mix as prepared above were added to each reaction test tube. The contents of the test tubes were then mixed by gentle pipetting. The tubes were then incubated in a PCR thermocycler for 20 min at 50°C.

The reaction was then stopped by adding 1 μl of 10X termination mix (0.1 M EDTA, 1 mg/ml glycogen).

Step B. Column Chromatography

The ¹²P-labeled cDNAs were separated from unincorporated ¹²P-labeled nucleotides and small (<0.1- kb) cDNA fragments using the following procedure for each test tube. A CHROMA SPIN-200 column (CLONTECH, Palo Alto, CA) was placed into a 1.5-ml microcentrifuge tube, the water was allowed to drain through the column by gravity flow until the surface of the gel beads emerged in the column matrix. The sample was then applied to the center of the gel bed's flat surface and allowed to be fully absorbed into the resin bed. 25 µl of ddH₂O were then applied and allowed to completely drain out of the column. 200 µl of ddH₂O were then applied and allowed to completely drain out of the column until there was no liquid left above the resin bed. The column was then transferred to a clean 1.5-ml microcentrifuge tube.

To collect the first fraction, 100 µl of ddH₂O were added to the column and allowed to completely drain out of the column. The second, third and fourth fractions were collected in analogous fashion. The tubes with fractions 1-4 were then placed in scintillation counter empty vials, and Cherenkov counts for each fraction were obtained in the tritium channel. The fractions which showed the highest Cerenkov counts were pooled.

Example 3 - Generation of Cy3-labeled hybridization polynucleotide target from polyA+RNA using postsynthesis labelling procedure

In this procedure for generating labeled cDNA target, polyA+RNA is first converted into cDNA that has primary amino groups which are subsequently coupled with Cy3 succinimide ester. This technology allows for a significant increase (about 10 fold) in activity of labeled polynucleotide target and therefore increases the overall sensitivity of detection of gene expression. The same procedure can be used for labeling two (or more) samples of RNA. In this case the cDNA synthesis step was the same for both samples but at the labeling step, each cDNA sample was labeled by different and distinguishable labels, e.g. Cy3 and Cy5, Alexa 532 and Bodipy TR, Fluorescein and tetramethyl rhodamine, etc. Each labeled probe was purified separately by column chromatography and, after normalization, were combined together in equal ratio and hybridized with a cDNA array. After hybridization, the detection procedure revealed both dye-labeled hybridized target simultaneously, based on the different spectral characteristics (emission wavelength) of the fluorescent labels.

A. cDNA synthesis

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The 10-μl reaction described below converted 1 μg of polyA+RNA into aminomodified first-strand cDNA.

For each cDNA synthesis reaction:

- 1. Enough master mix for all labeling reactions and 1 extra reaction was prepared to ensure sufficient volume.
- For each $10-\mu l$ labeling reaction, the following reagents were mixed:
 - $2 \mu l$ 5X First-strand buffer (250 μ M Tris-HC1 pH8.3; 375 mM KC1; 15 mM MgC12)
 - $1\,\mu l$ 10XdNTP mix (500 μ M dGTP, 500 μ M dCTP, 500 μ M dATP, 100 μ M dTTP,

and 100 µM allylamino dUTP)

- 1 μ l [α -12 P]dATP (Amersham, 3000 Ci/mmol, 10 mCi/ml)
- 3 μl H₂O
- 1 μl MMLV reverse transcriptase (Amersham, 200 units/ul)

5 ------ 8 μ] Final volume

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2. The following was combined in a 0.5-ml PCR test tube:

 $1 \mu g (1-2 \mu l)$ polyA+RNA sample

10 1 μ l 10x gene-specific primers mix (0.2 uM of each oligonucleotide ID No. 2,4,6,8,10,12,...... 1372) (from Table 1 of U.S. Patent Application No. 08/859,998, the disclosure of which is herein incorporated by reference.)

As a control in separate test tube 1 μ g of polyA+RNA sample was mixed with 1 μ l of oligo dT primer (SEQ ID NO. 1373 from Table 1 of U.S. Application No. 08/859,998).

- 3. ddH_2O was added to a final volume of 3 μ l.
- 4. The contents were mixed and the tubes were spun briefly in a microcentrifuge.
- 5. The tubes were incubated in preheated PCR thermocycler at 70°C for 2 min.
- 20 6. The temperature in the thermocycle was reduced down to 50°C and incubate for 2 min.
 - 7. $8 \mu l$ of master mix were added to each reaction test tube.
 - 8. The contents of the test tubes were mixed by gentle pipeting.
 - 9. The tubes were incubated in a PCR thermocycler for 30 min at 50°C.
- The reaction was stopped by increasing temperature up to 70°C for 5 min, then cooled to 37°C.
 - 11. 1 μ l of RNase H (10 units/ μ l) was added and the tubes were incubated at 37°C for 15 min.
 - 12. The reaction was stopped by adding 40 μ l of termination mix (0.3 M sodium acetate, pH 5.0, 1 mMEDTA).
 - 13. An equal volume (50 μ l) of phenol/chlorophorm/isoamyl alcohol mix (1: 1: 1/24 v/v) was added and extraction was performed. Phases were separated by centrifugation at 14,000 rpm for 10 min.

14. Upper water phase was collected and cDNA was precipitated by adding 2.5 volumes (about 120 μ l) of ethanol.

- 15. The precipitate was collected by centrifugation at 14,000 rpm for 10 min, the supernatant removed and the precipitate was washed with 80% ethanol.
- 5 16. The precipitate was air dried and dissolved in 10 μ l of 0. 1 M sodium bicarbonate buffer, pH 9.0.

Step B. Post synthesis labeling procedure.

- 1. I mg of Cy3 succinimide ester was dissolved in 10 μ l of dimethyl sulfoxide and 10 μ l of amino-modified cDNA generated at step 16 was added to it.
- 2. The mixture was incubated at room temperature overnight.

Step C. Column Chromatography

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To purify the Cy3-labeled cDNAs from the unconjugated label, the following was performed for each test tube:

- CHROMA SPIN-200 column (CLONTECH) was removed from refrigerator and allowed to warm at room temperature for about 1 hour. The column was inverted several times to completely resuspend the gel matrix. (Note: Check for air bubbles in the column matrix. If bubbles are visible, resuspend the matrix in the in the column buffer (ddH₂0) by inverting the column again).
- 2. The bottom cap from the column was removed, and then the top cap was slowly removed.
- 3. The column was placed into a 1.5-ml microcentrifuge tube.
- 4. The water was allowed to drain through the column by gravity flow until the surfaces
 25 of the gel beads in the column matrix were visible. (The top of the column matrix
 should be at the 0.75-ml mark on the wall of the column. If the column contains
 much less matrix, adjust the volume of the matrix to 0.75ml mark using matrix from
 another column.)
 - 5. The collected water was discarded.
- The sample was applied to the center of the gel bed's flat surface and allowed to be fully absorbed into the resin bed. Care was taken not allow any sample to flow along the inner wall of the column.

7. $25 \mu l$ of ddH₂0 were applied and allowed to completely drain out of the column.

- 8. Apply 200 μ l of ddH₂0 and allow the buffer to completely drain out of the column until there was no liquid left above the resin bed.
- 9. The column was transferred to a clean 1.5-ml microcentrifuge tube.
- $100 \mu l$ of ddH₂0 were added to the column and allowed to completely drain out of the column.
 - 11. The second, third and fourth fractions were collected by repeating steps 9-10.
 - 12. Cherenkov counts were obtained for each fraction by counting the entire sample in the tritium channel.
- 10 13. The fractions (usually fractions 2-3) which showed highest Cerenkov counts were pooled. Waste column and the fractions (usually fraction 1 and 4) which showed less than 10% counts from peak fractions.

Example 4 - Hybridization ¹²P-labeled cDNA Target with cDNA Array

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A solution of ExpressHybTM (CLONTECH) and sheared salmon testes DNA (Sigma) was prepared by prewarming 15 ml of ExpressHybTM at 50-60°C, heating 1.5 mg of sheared salmon testes DNA at 95-100°C for 5 min followed by chilling quickly on ice, and combining the resultant heat-denatured sheared salmon testes DNA with the prewarmed ExpressHybTM.

A cDNA Array as produced in Example 1 above was then placed in a hybridization bottle and 10 ml of the solution prepared above was added to the bottle. Prehybridization was performed for 30 min with continuous agitation at 72°C. Labeled cDNA probe (Example 1, about 200 ul, total about 2-5x10⁶ cpm) with 1/10th of the total volume (about 22 ul) of 10x denaturing solution (1 M NaOH, 10 mM EDTA) was mixed and incubated at 65°C for 20 min. 5 μl (1 μg/ul) of human Cot-1 DNA was then added, and an equal volume (about 225 μl) of 2x Neutralizing solution (1M NaHPO4, pH 7.0) was added and incubation continued at 65°C for 10 min. The mixtures were then combined and thoroughly mixed.

The prehybridization solution was replaced with the solution comprising the labeled oligonucleotide as prepared above and allowed to hybridize overnight with continuous agitation at 65°C. Following hybridization, the hybridization solution was carefully removed

and discarded, replaced with 200 ml of Wash Solution 1 (2X SSC, 1% SDS). The array was washed for 20 min with continuous agitation at 65°C. Washing was repeated four times.

Two additional 20-min washes were then performed in 200 ml of prewarmed Wash Solution 2 (0.1X SSC, 0.5% SDS) with continuous agitation at 65°C. Using forceps, the cDNA array was removed from the container and excess wash solution was removed by shaking.

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The damp membrane was immediately wrapped in plastic wrap, mounted on Whatman paper (3mm Chr) and exposed to x-ray film at -70°C with an intensifying screen.

Example 5 - Comparison Between Using Sets of Gene Specific Primers and oligo dT

³²P-labeled cDNA target were synthesized by M-MLV reverse transcriptase from a mixture 588 antisense gene-specific primers (B) or oligo dT(A) using placenta polyA+RNA as a template as described in Example 2. Primer extension products generated by reverse transcription were purified by gel filtration as described in Example 2 and hybridized separately with two cDNA arrays comprising 588 human genes under identical conditions as described in Example 4. Signals which can be detected by using cDNA target generated using the set of gene specific primers but can not be detected by using conventional target generated with oligo dT primers were observed. Note, the level of non-specific background detected as signal generated by membrane alone outside of the regions with immobilized probes generated by target generated using oligo dT primers was significantly higher in comparison with the background generated by the target generated by using the sets of gene specific primers.

Example 6 - Generation of cDNA array probe immobilized on glass slides.

50 cDNA fragments corresponding to 50 different human genes were amplified from plasmid clones containing corresponding cDNA fragments in 96 well plates using combination of vector primer ID No. 1376 and ID No. 1377 or sense and antisense genespecific primers: ID No. 1+2, 3+4,5+6,7+8,.... 100+101 (from Table 1 of U.S. Patent Application No. 08/859,998, the disclosure of which is herein incorporated by reference). Amplification was conducted in a 400-μl volume containing 2 ng of plasmid DNA, 40 mM Tricine-KOH (pH 9.2 at 22°C), 3.5 mM Mg(OAc), 10 MM KOAc, 75 μg/ml BSA, 200 μM

of each dATP, dGTP, dCTP and dTTP, 0.2 μ M of each primers and 2 μ l of KlenTaq Polymerase mix (CLONTECH). Temperature parameters of the PCR reactions were as follows: 1 min at 95°C followed by 30 cycles of 95°C for 15 sec and 68°C for 2 min; followed by a 10-min final extension at 68°C. PCR products were examined on 1.2% agarose/EtBr gels in 1 x TBE buffer. As a DNA size marker, a 1 Kb DNA Ladder was used. 5 ds cDNA was then precipitated by addition of a 10% volume of 3M sodium acetate (pH 5-0) (about 40 μ l) and 2.5 volumes of 96% ethanol (about 1 ml). After vortexing, the tube was immediately centrifuged at 14,000 r.p.m. in a microcentrifuge for 20 min. The pellet was washed with 80% ethanol without vortexing, centrifuged as above for 10 min, air dried, and dissolved in 10 μ l of deionized water. Yield of ds cDNA after amplification step was about 10 20 μg . The ds cDNA was solved in 10 μl of distilled water, 10 μl of 1 M sodium carbonate buffer, pH 9.5, was added and the ds cDNA was denaturated by heating at 94°C for 5 min and cooled down. The treated glass slides were prepared as following: Glass slides were cleaned overnight in 25% solution of nitric acid at room temperature, washed 3 times by acetone, treated with 1% aminopropyl-trimethoxysilane for 3 hrs at room temperature, 15 washed two times with acetone, heated at 120°C for 6 hrs and then treated with 0.2 % of para-phenylendiisothiocyanate (95:5 acetone-water solution) at room temperature for 3 hrs, then washed two times by acetone and dried in vacuum with desiccant. All cDNA probes were transferred in 384-well plate and printed on treated glass slides using 384 pin tool and Biomek 2000 (Beckman) robot. After printing, the arrays were incubated in wet chamber at 20 37°C overnight, then ultraviolet-cross linked to the surface by subjecting the slides to 30 mJ of energy (Stratagene Stratalinker). The arrays were washed with 1% of sodium borohydrate in 0.1 M NaOH, then washed 3 times in distilled water, dried in vacuum and stored with desiccant.

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Example 7- Hybridization Cy3 -labeled cDNA Target (or Cy3/Cy5 labeled cDNA targets) with glass cDNA array

- 1. A solution of ExpressHyb (CLONTECH) and sheared salmon testes DNA (Sigma) was prepared as follows:
 - a. 5 ml of ExpressHyb™ was prewarmed at 50-60°C.

b. 0.5 mg of the sheared salmon testes DNA was heated at 95-100 °C for 5 min, and then chilled quickly on ice.

- Heat-denatured sheared salmon testes DNA was mixed with prewarmed
 ExpressHyb.
- 5 2. The glass cDNA array was placed in a hybridization container, and 1 ml of the solution prepared in step 1 above was added.

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- 3. Prehybridization was conducted for 5 min with continuous agitation at 65°C.
- 4. Labeled cDNA probe as prepared in example 3, step C13, above, (about 200 μ l) was mixed with 2 μ l (1 μ g/ μ l) of human Cot- I DNA, and denaturated at 99°C for 2 min.
- 5. The mixture prepared in Step 4 was added to the hybridization box from Step 3 and the two solutions were mixed together thoroughly. The container was sealed by sealing tape.
- 6. Hybridization was allowed to proceed overnight with continuous agitation at 65°C.
- The hybridization solution was carefully removed and discarded in an appropriate container, and replaced with 10 ml of Wash Solution 1 (2X SSC, 0.1% SDS). The array was washed for 10 min with continuous agitation at 65°C. The step was repeated two times.
- 8. Additional 10-min washes were performed in 10 ml of Wash Solution 2 (0. 1 X SSC,
 20 0.1% SDS) with continuous agitation at 65°C.
 - Using forceps, the cDNA array was removed from the container, briefly washed in 0.
 1XSSC and excess buffer was removed from surface by centrifugation in a Beckman
 CS-6R centrifuge at 2000 rpm.
- Glass arrays were scanned using a custom-built laser scanner equipped by green (Cy3
 chanel) and red (Cy5 chanel) solid state laser built in UCLA. Images were scanned at a resolution of 20 μm per pixel.

It is evident from the above results and discussion that the subject invention provides a rapid, high throughput means to simply and quickly obtain a broad-scale screening of gene expression in a variety of different samples. Only simple hybridization protocols need be employed with the subject arrays, and signals can be detected using any convenient and readily available detection device. Despite their simplicity, assays conducted with the

subject arrays yield a large amount of information regarding the expression of numerous different and important genes in a particular sample at substantially the same time, and thus have use in many different types of applications, including drug discovery and characterization, disease research, and the like.

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All publications and patent applications cited in this specification are herein incorporated by reference as if each individual publication or patent application were specifically and individually indicated to be incorporated by reference. The citation of any publication is for its disclosure prior to the filing date and should not be construed as an admission that the present invention is not entitled to antedate such publication by virtue of prior invention.

Although the foregoing invention has been described in some detail by way of illustration and example for purposes of clarity of understanding, it is readily apparent to those of ordinary skill in the art in light of the teachings of this invention that certain changes and modifications may be made thereto without departing from the spirit or scope of the appended claims.

WHAT IS CLAIMED IS:

1. An array comprising a plurality of polynucleotide spots stably associated with the surface of a solid support, wherein a portion of said plurality of polynucleotide spots comprise a polynucleotide probe composition made up of unique polynucleotides and all of the unique polynucleotides on said array correspond to genes of a specific type.

- 2. The array according to Claim 1, wherein said polynucleotides of said array have an average length of from about 120 to 1000 nt.
- 10 3. The airay according to Claims 1 or 2, wherein each of said unique polynucleotides does not cross hybridize with the polynucleotides of any other polynucleotide probe composition on the array.
- 4. The array according to Claims 1, 2 or 3, wherein said polynucleotide probe composition comprises a population of single stranded identical polynucleotides.
 - 5. The array according to Claims 1, 2 or 3, wherein said polynucleotide probe composition comprises a population of two different complementary single stranded polynucleotides.

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- 6. The array according to any of the preceding claims, wherein the density of spots on said array does not exceed about 500/cm².
- 7. The array according to any of the preceding claims, wherein the number of spots on said array ranges from about 50 to 1000.
 - 8. The array according to any of the preceding claims, wherein said array is selected from the group consisting of a human array, a mouse array, a cancer array, an apoptosis array, a human stress array, an oncogene/tumor suppressor array, a cell-cell interaction array, a cytokine and cytokine receptor array, a rat array, a blood array, a mouse stress array, and a neuroarray.

9. The array according to any of the preceding claims, wherein said solid support is flexible.

- 10. The array according to any of the preceding claims, wherein said solid support is rigid.
 - 11. The array according to any of the preceding claims, wherein said polynucleotide probes of said array are those listed in a table selected from the group consisting of: Table 1, Table 2, Table 3, Table 4, Table 5, Table 6, Table 7 and Table 8.

12. A method of preparing an array according to any of the preceding claims, said method comprising:

enzymatically generating said unique polynucleotides; and stably associating said enzymatically-generated, complementary, unique polynucleotides on the surface of said solid support.

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- 13. A set of a representative number of distinct gene specific primers comprising gene specific primers corresponding to at least twenty distinct genes.
- 20 14. The set of gene specific primers according to Claim 13, wherein at least two of the twenty distinct genes are from different gene functional classes.
 - 15. The set of gene specific primers according to Claim 14, wherein the set comprises from 20 to 10,000 gene specific primers.
 - 16. The set of gene specific primers according to Claims 13, 14 or 15, wherein the set comprises primers capable of amplifying at least a portion of the polynucleotides present on an array according to any of Claims 1 to 11.
- 30 17. The set of gene specific primers according to Claim 16, wherein the set comprises primers capable of amplifying at least 20 of the polynucleotides present on an array according to any of Claims 1 to 11.

18. A method for detecting expression of a gene using a hybridization assay, said method comprising:

contacting at least one labeled target polynucleotide sample with an array according to any of Claims 1 to 11 under hybridization conditions sufficient to produce a hybridization pattern; and

detecting said hybridization pattern.

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- 19. The method according to Claim 18, wherein said method further comprises washing said array prior to said detecting step.
- 20. The method according to Claims 18 or 19, wherein said method further comprises preparing said labeled target polynucleotide sample.
- 21. The method according to Claim 20, wherein said preparation comprises:
 obtaining a sample of nucleic acids from a physiological source; and generating a population of labeled nucleic acids from the nucleic acids sample by using a set of a representative number of distinct gene specific primers according to any of Claims 13 to 17;

whereby said labeled target polynucleotide sample is produced.

- 22. The method according to Claims 20 or 21, wherein said preparing comprises conjugating a detectable label to a functionalized target polynucleotide.
- 23. The method according to any of Claims 18 to 22, where said method furthercomprises:

generating a second hybridization pattern; and comparing said hybridization patterns.

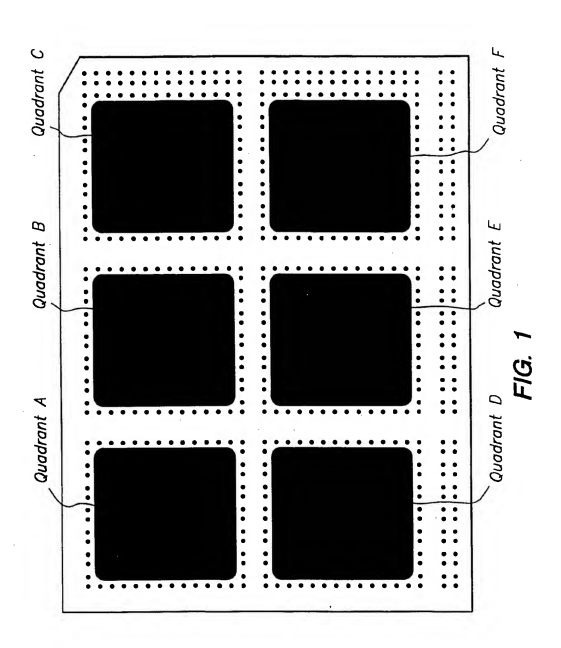
The method according to Claim 23, wherein said hybridization patterns are generatedon the same array.

25. The method according to Claim 23, wherein the second hybridization patters are generated on different arrays.

A kit for use in a hybridization assay, said kit comprising: an array according to any of Claims 1 to 11.

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- 27. The kit according to Claim 26, wherein said kit further comprises reagents for generating a labeled target polynucleotide sample.
- 10 28. The kit according to Claims 27, wherein said reagents comprise a set of a representational number of gene specific primers according to any of Claims 13 to 17.
 - 29. A kit for use in detecting the differential expression of genes of a plurality of physiological sources, the kit comprising:
- a set of a representative number of distinct gene specific primers according to any of Claims 13 to 17.



International application No. PCT/US98/10561

A. CLASSIFICATION OF SUBJECT MATTER IPC(6): C12Q 1/68; C12P 19/34; C07H 21/02, 21/04 US CL: 435/6, 91.1, 91.2; 536/23.1, 24.3, 24.31, 24.32, 24.33, 24.5 According to International Patent Classification (IPC) or to both national classification and IPC B. FIELDS SEARCHED Minimum documentation searched (classification system followed by classification symbols) U.S.: 435/6, 91.1, 91.2; 536/23.1, 24.3, 24.31, 24.32, 24.33, 24.5 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched						
Documentation restrict differ than imminute accommentation to the extent that rate accommend are more as a second and accommend						
Electronic data base consulted during the international search (name of data base and, where practicable, search terms used) Please See Extra Sheet.						
C. DOCUMENTS CONSIDERED TO BE RELEVANT						
Category* Citation of document, with indication, where a	opropriate, of the relevant passages Relevant to claim No.					
The in vitro acquisition of T cel	EHLERS et al. Differentiation of T cell lymphokine gene expression: The in vitro acquisition of T cell memory. J. Experimental Medicine. January 1991, Vol. 173, pages 25-36, see entire document.					
	CHALIFOUR et al. A method for analysis of gene expression patterns. Analytical Biochemistry. 1994, Vol. 216, pages 299-304, see entire document.					
Y ZHAO et al. High-density cDNA filt for large-scale, quantitative analysis of Vol. 156, pages 207-213, see entire of						
X Further documents are listed in the continuation of Box C. See patent family annex.						
* Special categories of cited documents: *A* document defining the general state of the art which is not considered to be of particular relevance *E* earlier document published on or after the international filing date *L* document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) *O* document referring to an oral disclosure, use, exhibition or other means *P* document published prior to the international filing date but later than the priority date claimed Date of the actual completion of the international search 24 JUNE 1998	later document published after the international filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention 'X' document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step when the document is taken alone 'Y' document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art '&' document member of the same patent family Date of mailing of the international search report					
Name and mailing address of the ISA/US Commissioner of Patents and Trademarks Box PCT Washington, D.C. 20231 Facsimile No. (703) 305-3230	Authorized offices JEFFREY FREDMAN Telephone No. (703) 308-0196					

International application No. PCT/US98/10561

C (Castinus	tion). DOCUMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where appropriate, of the relevant p	assages	Relevant to claim No
Y	NGUYEN et al. Differential gene expression in the murine thymus assayed by quantitative hybridization of arrayed cDNA clones. Genomics. 1995, Vol. 29, pages 207-216, see entire document.		1-3, 13-15
Y	Atlas human cDNA expression array I. Clontechniques. April 1997, pages 4-7, see entire document.		1-3, 13-15
Y	SCHENA et al. Parallel human genome analysis: Microarray-based expression monitoring of 1000 genes. Proc. Natl. Acad. Sci. October 1996, Vol. 93, pages 10614-10619, see entire document.		1-3, 13-15
Y	GOODWIN et al. Cloning of the human and murine interleukin 7 receptors: demonstration of a soluble, form and homology to a new receptor superfamily. Cell. 23 March 1990, Vol. 60, pages 941-951, see entire document.		1-3, 13-15
Y	LEONARD et al. Molecular cloning and expression of cDNAs for the human interleukin-2 receptor. Nature. 18 October 1984, Vol. 311, pages 626-631, see entire document.		1-3, 13-15
Y	GOODWIN et al. Human interleukin 7: Molecular cloning and growth factor activity on human and murine B-lineage cells. Proc. Natl. Acad. Sci. (USA). January 1989, Vol. 86, pages 302-306, see entire document.		1-3, 13-15
Y	NISHI et al. Cloning and expression of a novel variant of interferon gamma cDNA. J. Biochem. 1985, Vol. 97, No. 153-159, see entire document.		1-3, 13-15
	,		
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International application No. PCT/US98/10561

Box I Observations where certain claims were found unscarchable (Continuation of item 1 of first sheet)				
This international report has not been established in respect of certain claims under Article 17(2Xa) for the following reasons:				
1. Claims Nos.: because they relate to subject matter not required to be searched by this Authority, namely:				
2. Claims Nos.: because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:				
3. X Claims Nos.: 4-12, 16-19 because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).				
Box II Observations where unity of invention is lacking (Continuation of item 2 of first sheet)				
This International Searching Authority found multiple inventions in this international application, as follows:				
Please See Extra Sheet.				
1. As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.				
2. As all searchable claims could be searched without effort justifying an additional fee, this Authority did not invite paymen of any additional fee.				
As only some of the required additional search fees were timely paid by the applicant, this international search report covers only those claims for which fees were paid, specifically claims Nos.:				
4. X No required additional search fees were timely paid by the applicant. Consequently, this international search report i restricted to the invention first mentioned in the claims; it is covered by claims Nos.: 1-29, species of SEQ ID NOs: 1-10				
Remark on Protest The additional search fees were accompanied by the applicant's protest.				
No protest accompanied the payment of additional search fees.				

International application No. PCT/US98/10561

B. FIELDS SEARCHED

Electronic data bases consulted (Name of data base and where practicable terms used):

APS, MEDLINE, BIOSIS, CAPLUS

search terms: array, support, bead, nitrocellulose, nylon, filter, hybridize, anneal, DNA, RNA, gene, nucleic, oligo, polynucleotide, spot, pattem, primer

BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING

This ISA found multiple inventions as follows:

This application contains claims directed to more than one species of the generic invention. These species are deemed to lack Unity of Invention because they are not so linked as to form a single inventive concept under PCT Rule 13.1. In order for more than one species to be searched, the appropriate additional search fees must be paid. The species are as follows:

Each of the sequences found in Tables 1-8.

The species listed above do not relate to a single inventive concept under PCT Rule 13.1 because, under PCT Rule 13.2, the species lack the same or corresponding special technical features for the following reasons:

Each of the sequences found in Tables 1-8 represents a different nucleic acid species which are not joined by a corresponding technical feature such as encoding a similar protein.

According the Official Gazette Notice in October 1996, "Under the Unity of Invention Standard in an International Application or National Stage Application Filed Under 35 U.S.C. § 371, Up to Ten Nucleotide Sequences Will Be Searched and/or Examined Without Payment of An Additional Fee".